(19) World Intellectual Property Organization International Bureau



(43) International Publication Date 27 June 2002 (27.06.2002)

PCT

(10) International Publication Number WO 02/50096 A1

- (51) International Patent Classification⁷: C07H 21/04, C07K 14/435, 16/18, C12N 15/12, 15/63, A61K 38/17, 39/395, 31/7105, 48/00, A61P 25/08
- (21) International Application Number: PCT/AU01/01648
- (22) International Filing Date:

20 December 2001 (20.12.2001)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

PR 2203

20 December 2000 (20.12.2000) A

- (71) Applicant (for all designated States except US): BIONOMICS LIMITED [AU/AU]; 31 Dalgleish Street, Thebarton, S.A. 5031 (AU).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): WALLACE, Robyn, Heather [AU/AU]; 51 Auricchio Avenue, St Marys S.A. 5042 (AU). MULLEY, John, Charles [AU/AU]; 13 Dunkley Avenue, Firle, S.A. 5046 (AU). BERKOVIC, Samuel,

Frank [AU/AU]; 7 Polo Parade, Caulfield North, VIC 3161 (AU).

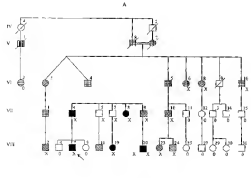
- (74) Agent: GRIFFITH HACK; GPO Box 3125, Brisbane, QLD 4001 (AU).
- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

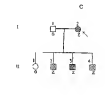
Published:

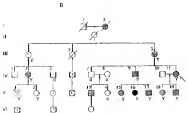
with international search report

[Continued on next page]

(54) Title: SODIUM-CHANNEL ALPHA1-SUBUNIT AND THEIR POLYPEPTIDES AND THEIR TREATMENT OF GENERALISED EPILEPSY WITH FEBRILE SEIZURES PLUS









02/50096 A

(57) Abstract: The mutations D188V, V1353L, I1656M in the neuronal gene sodium-channel alpha1-subunit, SCN1A, are disclosed. The methods of using their associated polypeptides for treating sodium channel dysfunction disorders including generalised epilepsy are also disclosed.

WO 02/50096 A1



For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

- 1 -

Mutations in Neuronal gene sodium-channel alpha1-subunit and their polypeptides and their treatment of generalised epilepsy with febrile seizures plus.

Technical Field

5

10

15

20

25

30

The present invention relates to mutations in the alpha subunit of mammalian voltage-gated sodium channels which are associated with idiopathic epilepsies and other disorders such as malignant hyperthermia, myasthenia, episodic ataxia, neuropathic and inflammatory pain, Alzheimer's disease, Parkinson's disease, schizophrenia, hyperekplexia, myotonias and cardiac arrhythmias, and to polymorphisms in the gene encoding the alpha subunit.

Background Art

epilepsy with Generalised febrile seizures plus (GEFS+; MIM 604236) was first described by Scheffer and Berkovic (1997) and is now recognised as a common epilepsy syndrome (Singh et al. 1999; Baulac et al. 1999; Moulard et al. 1999; Peiffer et al. 1999; Scheffer et al. 2000). Although GEFS+ is familial, it was initially difficult to recognise it as a distinct syndrome, because of clinical heterogeneity within each family. The common phenotypes are typical febrile seizures (FS) and febrile seizures plus (FS+); FS+ differs from FS in that the attacks with fever continue beyond age 6 years and/or include afebrile tonic-clonic seizures. Less common phenotypes include FS+ associated with absences, myoclonic or atonic seizures, and even more-severe syndromes such as myoclonic-astatic phenotypic epilepsy. That such diversity associated with the segregation of a mutation in a single gene was established with the identification of a mutation in the voltage gated sodium channel beta-1 subunit gene (Wallace et al. 1998). This mutation (C121W) (SCN1B) conserved cysteine residue, disrupting changes putative disulfide bridge, which results in in vitro loss of function of the beta-1 subunit. Without a functional beta-1 subunit the rate of inactivation of sodium channel alpha subunits decreases, which may cause increased sodium

- 2 -

influx, resulting in a more depolarised membrane potential and hyperexcitability. Modifier genes or the environment may interact with the SCN1B gene to account for clinical heterogeneity, but the rarity of SCN1B mutations (Wallace et al. 1998) strongly suggested additional genes of large effect underlie GEFS+ in other families (Singh et al. 1999).

GEFS+ in four families has been mapped to chromosome 2q (Baulac et al. 1999; Moulard et al. 1999; Peiffer et al. 1999; Lopes-Cendes et al. 2000). Recently, mutations in the neuronal voltage gated sodium channel alpha-1 (SCN1A) subunit were described in two GEFS+ families (Escayg et al. 2000). The mutations (T875M and R1648H) are located in highly conserved S4 transmembrane segments of the channel which are known to have a role in channel gating. It was suggested that these mutations may reduce the rate of inactivation of SCN1A and therefore have a similar effect as the beta-1 subunit mutation.

GEFS+ is clearly a common complex disorder, with a strong genetic basis, incomplete penetrance and genetic and phenotypic heterogeneity. Febrile seizures occur in 3% of the population, and thus this phenotype may occur sporadically in GEFS+ families, in addition to occurring as a result of an inherited mutation in the GEFS+ gene al 1998). Also, although some families (Wallace et segregate an autosomal dominant gene of major effect, in many cases clinical genetic evidence, such as bilineality, suggests that for some small families the disorder is multifactorial (Singh et al 1999). Despite this, large families continue to be ascertained and with critical provide opportunities phenotypic analysis, they localise and ultimately identify the genes involved.

Disclosure of the Invention

10

15

20

25

30

35 The present inventors have identified three new mutations in the alpha-1 subunit (SCN1A) of the voltage-gated sodium channel that are associated with epilepsy, in

particular generalized epilepsy with febrile seizures plus (GEFS+), and also determined the nucleotide sequence in that gene.

- 3 -

According to one aspect of the present invention there is provided an isolated DNA molecule encoding a mutant alpha subunit of a mammalian voltage-gated sodium channel, wherein a mutation event selected from the group consisting of point mutations, deletions, insertions and rearrangements has occurred and said mutation event disrupts the functioning of an assembled sodium channel so as to produce an epilepsy phenotype, with the proviso that the mutation event is not a C2624T transition or a G4943A transition in an alpha-1 subunit.

10

15

20

25

30

35

Preferably said mutation event is a point mutation.

Typically the mutation event occurs in an intracellular loop, preferably in the intracellular loop between transmembrane segments 2 and 3 of domain I, in the S4 segment of domain IV at amino acid position 1656, or in an S5 segment of a transmembrane domain. Preferably the mutation creates a phenotype of generalised epilepsy with febrile seizures plus.

In one form of the invention the mutation is in exon 4 of SCN1A and results in replacement of a highly conserved aspartic acid residue with a valine residue at amino acid position 188. The D188V mutation lies in the intracellular loop just outside the S3 segment of domain I of SCN1A and occurs as a result of an A to T nucleotide substitution at position 563 of the SCN1A coding sequence as shown in SEO ID NO:1.

In a further form of the invention the mutation is in exon 21 of SCN1A and results in the replacement of a highly conserved valine residue with a leucine residue at amino acid position 1353. The V1353L mutation is located in the S5 segment of domain III of SCN1A and occurs as a result of a G to C nucleotide substitution at position 4057 of the SCN1A coding sequence as shown in SEQ ID NO:3.

In a still further form of the invention the mutation

10

15

20

25

30

35

- 4 -

is in exon 26 of SCN1A and results in the replacement of a highly conserved isoleucine residue with a methionine residue at amino acid position 1656. The I1656M mutation is located in the S4 segment of domain IV of SCN1A and occurs as a result of a C to G nucleotide substitution at position 4968 of the SCN1A coding sequence as shown in SEQ ID NO:5.

The nucleotide sequence of the gene set forth in SEQ ID NO:89 also forms a part of the invention. In addition, the polymorphisms identified in Table 3 form part of the invention (SEQ ID Numbers:7-9 and 11).

The present invention also encompasses DNA molecules in which one or more additional mutation events selected from the group consisting of point mutations, deletions, insertions and rearrangements have occurred. Any such DNA molecule will have the mutation associated with epilepsy described above and will be functional, but otherwise may vary significantly from the DNA molecules set forth in SEQ ID NO:1, 3 and 5.

The nucleotide sequences of the present invention can be engineered using methods accepted in the art for a variety of purposes. These include, but are not limited to, modification of the cloning, processing, and/or expression of the gene product. PCR reassembly of gene fragments and the use of synthetic oligonucleotides allow the engineering of the nucleotide sequences of the present invention. For example, oligonucleotide-mediated site-directed mutagenesis can introduce further mutations that create new restriction sites, alter expression patterns and produce splice variants etc.

As a result of the degeneracy of the genetic code, a number of polynucleotide sequences, some that may have minimal similarity to the polynucleotide sequences of any known and naturally occurring gene, may be produced. Thus, the invention includes each and every possible variation of a polynucleotide sequence that could be made by selecting combinations based on possible codon choices.

10

15

20

25

30

35

- 5 -

These combinations are made in accordance with the standard triplet genetic code as applied to the polynucleotide sequences of the present invention, and all such variations are to be considered as being specifically disclosed.

The DNA molecules of this invention include cDNA, genomic DNA, synthetic forms, and mixed polymers, both sense and antisense strands, and may be chemically or biochemically modified, or may contain non-natural or derivatised nucleotide bases as will be appreciated by those skilled in the art. Such modifications include labels. methylation, intercalators, alkylators modified linkages. In some instances it may be advantageous to produce nucleotide sequences possessing a substantially different codon usage than that polynucleotide sequences of the present invention. For example, codons may be selected to increase the rate of expression of the peptide in a particular prokaryotic or eukaryotic host corresponding with the frequency that particular codons are utilized by the host. Other reasons to alter the nucleotide sequence without altering the encoded amino acid sequences include the production of RNA transcripts having more desirable properties, such as a greater half-life, than transcripts produced from the naturally occurring mutated sequence.

The invention also encompasses production of DNA sequences of the present invention entirely by synthetic chemistry. Synthetic sequences may be inserted into expression vectors and cell systems that contain the necessary elements for transcriptional and translational control of the inserted coding sequence in a suitable host. These elements may include regulatory sequences, promoters, 5' and 3' untranslated regions and specific initiation signals (such as an ATG initiation codon and Kozak consensus sequence) which allow more efficient translation of sequences encoding the polypeptides of the present invention. In cases where the complete coding

- 6 -

sequence, including the initiation codon and upstream regulatory sequences, are inserted into the appropriate expression vector, additional control signals may not be needed. However, in cases where only coding sequence, or a fragment thereof, is inserted, exogenous translational control signals as described above should be provided by the vector. Such signals may be of various origins, both natural and synthetic. The efficiency of expression may be enhanced by the inclusion of enhancers appropriate for the particular host cell system used (Scharf et al., 1994).

The invention also includes nucleic acid molecules that are the complements of the sequences described herein.

10

15

20

25

30

35

According to still another aspect of the present invention there is provided an isolated DNA molecule consisting of the nucleotide sequence set forth in any one of SEQ ID NOS:1, 3, 5, 7, 8, 9, 11 and 89.

The present invention allows for the preparation of purified polypeptides or proteins from the polynucleotides of the present invention, or variants thereof. In order to do this, host cells may be transformed with a DNA molecule described above. Typically said host cells are transfected with an expression vector comprising a DNA molecule according to the invention. A variety expression vector/host systems may be utilized to contain express sequences encoding polypeptides invention. These include, but are not limited to, microorganisms such as bacteria transformed with plasmid or cosmid DNA expression vectors; yeast transformed with yeast expression vectors; insect cell systems infected with viral expression vectors (e.g., baculovirus); or mouse or other animal or human tissue cell systems. Mammalian cells can be used to express a protein using various expression vectors including plasmid, cosmid and viral systems such as a vaccinia virus expression system. The invention is not limited by the host cell employed.

The polynucleotide sequences, or variants thereof, of

10

15

20

25

30

35

- 7 -

the present invention can be stably expressed in cell lines to allow long term production of recombinant proteins in mammalian systems. Sequences encoding the polypeptides of the present invention can be transformed into cell lines using expression vectors which may contain viral origins of replication and/or endogenous expression elements and a selectable marker gene on the same or on a separate vector. The selectable marker confers resistance to a selective agent, and its presence allows growth and of which successfully express recovery cells introduced sequences. Resistant clones of stably transformed cells may be propagated using tissue culture techniques appropriate to the cell type.

The protein produced by a transformed cell may be secreted or retained intracellularly depending on the sequence and/or the vector used. As will be understood by those of skill in the art, expression vectors containing polynucleotides which encode a protein may be designed to contain signal sequences which direct secretion of the protein through a prokaryotic or eukaryotic cell membrane.

In addition, a host cell strain may be chosen for its ability to modulate expression of the inserted sequences or to process the expressed protein in the desired fashion. Such modifications of the polypeptide include, are not limited to, acetylation, glycosylation, phosphorylation, and acylation. Post-translational cleavage of a "prepro" form of the protein may also be specify protein targeting, folding, activity. Different host cells having specific cellular characteristic mechanisms machinery and for translational activities (e.g., CHO or HeLa cells), are available from the American Type Culture Collection (ATCC) and may be chosen to ensure the correct modification and processing of the foreign protein.

When large quantities of the gene are needed, such as for antibody production, vectors which direct high levels of expression of this protein may be used, such as those

- 8 -

containing the T5 or T7 inducible bacteriophage promoter. The present invention also includes the use of the expression systems described above in generating and isolating fusion proteins which contain important functional domains of the protein. These fusion proteins are used for binding, structural and functional studies as well as for the generation of appropriate antibodies.

In order to express and purify the protein as a fusion protein, the appropriate polynucleotide sequences of the present invention are inserted into a vector which contains a nucleotide sequence encoding another peptide (for example, glutathionine-s-transferase). The fusion protein is expressed and recovered from prokaryotic or eukaryotic cells. The fusion protein can then be purified by affinity chromatography based upon the fusion vector sequence. The desired protein is then obtained by enzymatic cleavage of the fusion protein.

10

15

20

25

30

35

Fragments of polypeptides of the present invention may also be produced by direct peptide synthesis using solid-phase techniques. Automated synthesis may be achieved by using the ABI 431A Peptide Synthesizer (Perkin-Elmer). Various fragments of this protein may be synthesized separately and then combined to produce the full length molecule.

According to still another aspect of the present invention there is provided an isolated polypeptide, said polypeptide being a mutant alpha subunit of a mammalian voltage-gated sodium channel, wherein a mutation event selected from the group consisting of point mutations, deletions, insertions and rearrangements has occurred and said mutation event disrupts the functioning of an assembled sodium channel so as to produce an epilepsy phenotype, with the proviso that said mutation event is not a T875M transition or a R1648H transition in an alpha-1 subunit.

Preferably said mutation event occurs in an intracellular loop, preferably in the intracellular loop

5

10

15

20

25

30

35

- 9 -

between transmembrane segments 2 and 3 in domain I, in the S4 segment of domain IV at amino acid position 1656, or in an S5 segment of a transmembrane domain of SCN1A. Preferably the mutation creates a phenotype of generalised epilepsy with febrile seizures plus.

In one form of the invention the mutation event is a substitution in which a highly conserved aspartic acid residue is replaced with a valine residue located in the intracellular domain located just outside the S3 segment of domain I of SCN1A. Preferably the substitution is a D188V transition as illustrated in SEO ID NO:2.

In a further form of the invention the mutation event is a substitution in which a highly conserved valine residue is replaced with a leucine residue located in the S5 segment of domain III of SCN1A. Preferably the substitution is a V1353L transition as illustrated in SEQ ID NO:4.

In a still further form of the invention the mutation event is a substitution in which a highly conserved isoleucine residue is replaced with a methionine residue located in the S4 segment of domain IV of SCN1A. Preferably the substitution is a I1656M transition as illustrated in SEQ ID NO:6.

In addition, the polymorphisms identified in Table 3 form part of the invention (SEQ ID Numbers:10 and 12). These polymorphisms may reflect changes in SCN1A which result in subtle changes of function of the sodium channel. These subtle changes may predispose individuals to epilepsy and when expressed in combination with other ion channel changes may lead to specific sub-types of the disease (see PCT/AU01/00872).

The isolated polypeptides of the present invention may have been subjected to one or more mutation events selected from the group consisting of substitutions, deletions, insertions and rearrangements in addition to the mutation associated with epilepsy. Typically these mutation events are conservative substitutions.

5

10

15

20

25

30

3.5

- 10 -

According to still another aspect of the present invention there is provided an isolated polypeptide comprising the sequence set forth in any one of SEQ ID NO:2, 4, 6, 10 and 12.

According to still another aspect of the present invention there is provided a polypeptide consisting of the amino acid sequence set forth in any one of SEQ ID NO:2, 4, 6, 10 and 12.

According to still another aspect of the present there is provided an isolated polypeptide complex, said polypeptide complex being an assembled mammalian voltage-gated sodium channel, wherein a mutation event selected from the group consisting of substitutions, deletions, insertions and rearrangements has occurred in the alpha subunit of the complex. Mutations include those in the intracellular loop between transmembrane segments 2 and 3, the S4 segment of domain IV at amino acid position 1656, or in an S5 segment of a transmembrane domain of the In a particular aspect subunit. an assembled mammalian voltage-gated sodium channel bearing any such mutation in the alpha subunit will produce a phenotype of epilepsy, in particular generalised epilepsy with febrile seizures plus, or other disorders associated with sodium channel dysfunction including, but not restricted to, episodic malignant hyperthermia, myasthenia, neuropathic and inflammatory pain, Alzheimer's disease, Parkinson's disease, schizophrenia, hyperekplexia, hypo- and hyperkalaemic myotonias such as paralysis, paramyotonia congenita and potassium aggravated myotonia as well as cardiac arrhythmias such as long QT syndrome.

In a particular aspect there is provided a complex, being an assembled mammalian voltage-gated sodium channel, bearing a mutation in the intracellular loop between transmembrane segments 2 and 3, the S4 segment of domain IV at amino acid position 1656, or in an S5 segment of a transmembrane domain of the SCN1A subunit of the channel.

WO 02/50096

5

15

20

25

30

According to still another aspect of the present invention there is provided a method of preparing a polypeptide, said polypeptide being a mutant alpha subunit of a mammalian voltage-gated sodium channel, comprising the steps of:

- 11 -

PCT/AU01/01648

- (1) culturing host cells transfected with an expression vector comprising a nucleic acid molecule as described above under conditions effective for polypeptide production; and
- 10 (2) harvesting the mutant alpha subunit.

The mutant alpha subunit may also be allowed to assemble with other subunits of the sodium channel, whereby the assembled mutant sodium channel is harvested.

According to still another aspect of the invention there is provided a polypeptide which is the product of the process described above.

Substantially purified protein or fragments thereof can then be used in further biochemical analyses to establish secondary and tertiary structure for example by X-ray crystallography of crystals of the proteins or by nuclear magnetic resonance (NMR). Determination of the rational allows for design of structure interact with the mutated sodium pharmaceuticals to channel, alter the overall sodium channel protein charge configuration or charge interaction with other proteins, or to alter its function in the cell.

It will be appreciated that, having identified mutations involved in epilepsy in these proteins, the mutant sodium channel alpha subunits will be useful in applications which include a variety of further hybridisation and immunological assays to screen for and detect the presence of either a normal or mutated gene or gene product. The invention also enables therapeutic methods for the treatment of epilepsy and enables methods for the diagnosis of epilepsy with both wild-type and mutant nucleic acid molecules. In particular the invention enables treatment and diagnosis of generalised epilepsy

with febrile seizures plus, as well as other disorders associated with sodium channel dysfunction, as mentioned

- 12 -

5 Therapeutic Applications

above.

10

15

20

25

30

According to one aspect of the invention there is provided a method of treating epilepsy, in particular generalised epilepsy with febrile seizures plus, as well disorders associated with sodium other dysfunction, including but not restricted to, malignant hyperthermia, myasthenia, episodic ataxia, neuropathic and disease, pain, Alzheimer's inflammatory Parkinson's disease, schizophrenia, hyperekplexia, myotonias such as hypo- and hyperkalaemic periodic paralysis, paramyotonia congenita and potassium aggravated myotonia as well as cardiac arrhythmias such as long QT syndrome, comprising administering a selective antagonist, agonist or modulator of the sodium channel when a mutation event as described above has occurred, in particular, when it contains a mutation in the intracellular loop between transmembrane segments 2 and 3, in the S4 segment of domain IV at amino acid position 1656, or in an S5 segment of a transmembrane domain of an alpha subunit.

In still another aspect of the invention there is provided the use of a selective antagonist, agonist or modulator of the sodium channel when a mutation event as described above has occurred, in particular, to a sodium channel when it contains a mutation in the intracellular loop between transmembrane segments 2 and 3, in the S4 segment of domain IV at amino acid position 1656, or in an S5 segment of a transmembrane domain of an alpha subunit, said mutation being causative of a disorder including epilepsy, in particular generalised epilepsy with febrile seizures plus as well as other disorders associated with sodium channel dysfunction, including but not restricted to, malignant hyperthermia, myasthenia, episodic ataxia, neuropathic and inflammatory pain, Alzheimer's disease,

10

15

20

25

30

- 13 -

Parkinson's disease, schizophrenia, hyperekplexia, myotonias such as hypo- and hyperkalaemic periodic paralysis, paramyotonia congenita and potassium aggravated myotonia as well as cardiac arrhythmias such as long QT syndrome, in the manufacture of a medicament for the treatment of the disorder.

In one aspect of the invention a suitable antagonist or modulator will restore wild-type function to the sodium channels that contain a mutation in an alpha subunit including those that form part of this invention.

Using methods well known in the art, a mutant sodium channel may be used to produce antibodies specific for the mutant channel that is causative of the disease or to screen libraries of pharmaceutical agents to identify those that specifically bind the mutant sodium channel.

In one aspect, an antibody, which specifically binds to a mutant sodium channel, may be used directly as an antagonist or modulator, or indirectly as a targeting or delivery mechanism for bringing a pharmaceutical agent to cells or tissues that express the mutant sodium channel.

In a still further aspect of the invention there is provided an antibody which is immunologically reactive with a polypeptide as described above, but not with a wild-type sodium channel or subunit thereof.

In particular, there is provided an antibody to an assembled sodium channel containing a mutation causative of a disorder as described above, in a subunit comprising the receptor. Such antibodies may include, but are not limited to, polyclonal, monoclonal, chimeric, and single chain antibodies as would be understood by the person skilled in the art.

For the production of antibodies, various hosts including rabbits, rats, goats, mice, humans, and others may be immunized by injection with a polypeptide as described or with any fragment or oligopeptide thereof which has immunogenic properties. Various adjuvants may be used to increase immunological response and include, but

5

10

15

20

25

30

35

- 14 -

are not limited to, Freund's, mineral gels such as aluminum hydroxide, and surface-active substances such as lysolecithin. Adjuvants used in humans include BCG (bacilli Calmette-Guerin) and Corynebacterium parvum.

It is preferred that the oligopeptides, peptides, or fragments used to induce antibodies to the mutant sodium channel have an amino acid sequence consisting of at least 5 amino acids, and, more preferably, of at least 10 amino acids. It is also preferable that these oligopeptides, peptides, or fragments are identical to a portion of the amino acid sequence of the natural protein and contain the entire amino acid sequence of a small, naturally occurring molecule. Short stretches of sodium channel amino acids may be fused with those of another protein, such as KLH, and antibodies to the chimeric molecule may be produced.

Monoclonal antibodies to a mutant sodium channel may be prepared using any technique which provides for the production of antibody molecules by continuous cell lines in culture. These include, but are not limited to, the hybridoma technique, the human B-cell hybridoma technique, and the EBV-hybridoma technique. (For example, see Kohler et al., 1975; Kozbor et al., 1985; Cote et al., 1983; Cole et al., 1984).

Antibodies may also be produced by inducing in vivo production in the lymphocyte population or by screening immunoglobulin libraries or panels of highly specific binding reagents as disclosed in the literature. (For example, see Orlandi et al., 1989; Winter et al., 1991).

Antibody fragments which contain specific binding sites for a mutant sodium channel may also be generated. For example, such fragments include, F(ab')2 fragments produced by pepsin digestion of the antibody molecule and Fab fragments generated by reducing the disulfide bridges of the F(ab')2 fragments. Alternatively, Fab expression libraries may be constructed to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity. (For example, see Huse et al., 1989).

10

15

20

25

30

35

- 15 -

Various immunoassays may be used for screening to identify antibodies having the desired specificity. protocols for competitive binding Numerous orimmunoradiometric assays using either polyclonal monoclonal antibodies with established specificities are well known in the art. Such immunoassays typically involve the measurement of complex formation between a sodium channel and its specific antibody. A two-site, monoclonalbased immunoassay utilizing antibodies reactive to two non-interfering sodium channel epitopes is preferred, but a competitive binding assay may also be employed.

In a further aspect of the invention there provided a method of treating epilepsy, in particular generalised epilepsy with febrile seizures plus, as well other disorders associated with sodium dysfunction, including but not restricted to, malignant hyperthermia, myasthenia, episodic ataxia, neuropathic and Alzheimer's inflammatory pain, disease, Parkinson's disease, schizophrenia, hyperekplexia, myotonias such as hypo- and hyperkalaemic periodic paralysis, paramyotonia congenita and potassium aggravated myotonia as well as cardiac arrhythmias such as long QT syndrome, comprising administering an isolated DNA molecule which complement (antisense) of any one of the DNA molecules described above and which encodes an RNA molecule that hybridizes with the mRNA encoding a mutant sodium channel alpha subunit, to a subject in need of such treatment.

Typically, a vector expressing the complement of the polynucleotides of the invention may be administered to a subject in need of such treatment. Antisense strategies may use a variety of approaches including the use of antisense oligonucleotides, injection of antisense RNA, ribozymes, DNAzymes and transfection of antisense RNA expression vectors. Many methods for introducing vectors into cells or tissues are available and equally suitable for use in vivo, in vitro, and ex vivo. For ex vivo therapy, vectors may be introduced into stem cells taken

10

15

20

25

30

35

- 16 -

from the patient and clonally propagated for autologous transplant back into that same patient. Delivery by transfection, by liposome injections, or by polycationic amino polymers may be achieved using methods which are well known in the art. (For example, see Goldman et al., 1997).

In a still further aspect of the invention there is provided the use of an isolated DNA molecule which is the complement of a DNA molecule of the invention and which encodes an RNA molecule that hybridizes with the mRNA encoding a mutant sodium channel alpha subunit, in the manufacture of a medicament for the treatment of epilepsy, in particular generalised epilepsy with febrile seizures plus, as well as other disorders associated with sodium channel dysfunction, including but not restricted to, hyperthermia, myasthenia, episodic malignant neuropathic and inflammatory pain, Alzheimer's disease, schizophrenia, Parkinson's disease, hyperekplexia, hypoand hyperkalaemic myotonias such as periodic paralysis, paramyotonia congenita and potassium aggravated myotonia as well as cardiac arrhythmias such as long QT syndrome.

In a further aspect, a suitable agonist or modulator may include a small molecule that can restore wild-type activity of the sodium channel containing mutations in the alpha subunit as described above, or may include an antibody to a mutant sodium channel that is able to restore channel function to a normal level.

Small molecules suitable for therapeutic applications may be identified using nucleic acids and peptides of the invention in drug screening applications as described below.

In further embodiments, any of the agonists, antagonists, modulators, antibodies, complementary sequences or vectors of the invention may be administered alone or in combination with other appropriate therapeutic agents. Selection of the appropriate agents may be made by

- 17 -

PCT/AU01/01648

those skilled in the art, according to conventional pharmaceutical principles. The combination of therapeutic agents may act synergistically to effect the treatment or prevention of the various disorders described above. Using this approach, therapeutic efficacy with lower dosages of each agent may be possible, thus reducing the potential for adverse side effects.

Drug screening

WO 02/50096

10

15

20

25

30

3.5

According to still another aspect of the invention, peptides of the invention, particularly purified mutant sodium channel alpha subunit polypeptide and expressing these, are useful for the screening in candidate pharmaceutical agents a variety of techniques. It will be appreciated that therapeutic agents useful in the treatment of epilepsy, in particular generalised epilepsy with febrile seizures plus, as well other disorders associated with sodium as dysfunction, including but not restricted to, malignant hyperthermia, myasthenia, episodic ataxia, neuropathic and Alzheimer's disease, inflammatory pain, Parkinson's disease, schizophrenia, hyperekplexia, myotonias such as hypo- and hyperkalaemic periodic paralysis, paramyotonia congenita and potassium aggravated myotonia as well as cardiac arrhythmias such as long QT syndrome, are likely to show binding affinity to the polypeptides of the invention.

Such techniques include, but are not limited to, utilising eukaryotic or prokaryotic host cells that are stably transformed with recombinant molecules expressing the polypeptide or fragment, preferably in competitive binding assays. Binding assays will measure the formation of complexes between a mutated sodium channel alpha subunit polypeptide or fragment and the agent being tested, or will measure the degree to which an agent being tested will interfere with the formation of a complex between a mutated sodium channel alpha subunit polypeptide or fragment and a known ligand.

10

15

20

25

30

35

- 18 -

Another technique for drug screening provides highthroughput screening for compounds having suitable binding affinity to the mutant sodium channel alpha polypeptides or sodium channels containing these (see PCT published application WO84/03564). In this technique, large numbers of small peptide test compounds can be synthesised on a solid substrate and can be assayed through mutant sodium channel or mutant sodium channel alpha subunit polypeptide binding and washing. mutant sodium channel or mutant sodium channel alpha subunit polypeptide is then detected by methods well known in the art. In a variation of this technique, purified polypeptides of the invention can be coated directly onto plates to identify interacting test compounds.

The invention also contemplates the use of competition drug screening assays in which neutralizing antibodies capable of specifically binding the mutant sodium channel compete with a test compound for binding thereto. In this manner, the antibodies can be used to detect the presence of any peptide that shares one or more antigenic determinants of the mutant sodium channel.

The invention is particularly useful for screening compounds by using the polypeptides of the invention in transformed cells, transfected or injected oocytes, modelsbearing mutated sodium channel animal subunits (particularly those of the invention) such as transgenic animals or gene targeted (knock-in) animals (see below). A particular drug is added to the cells in culture or administered to an animal model containing a mutant sodium channel alpha subunit and the effect on the current of the channel is compared to the current of a cell or animal containing the wild-type sodium channel. Drug candidates that alter the current to a more normal level are useful for treating or preventing epilepsy, in particular generalised epilepsy with febrile seizures plus as well as other disorders associated with sodium channel dysfunction, as described above.

10

15

20

25

30

- 19 -

The polypeptides of the present invention may also be used for screening compounds developed as a result of combinatorial library technology. This provides a way to test a large number of different substances for their ability to modulate activity of a polypeptide. The use of peptide libraries is preferred (see WO 97/02048) with such libraries and their use known in the art.

A substance identified as a modulator of polypeptide function may be peptide or non-peptide in nature. Nonpeptide "small molecules" are often preferred for many in vivo pharmaceutical applications. In addition, a mimic or of substance mimetic the may be designed for pharmaceutical use. The design of mimetics based on a known pharmaceutically active compound ("lead" compound) common approach to the development of is pharmaceuticals. This often desirable where original active compound is difficult or expensive to synthesise or where it provides an unsuitable method of administration. In the design of a mimetic, particular parts of the original active compound that are important in determining the target property are identified. These parts or residues constituting the active region of the compound are known as its pharmacophore. Once found, the pharmacophore structure is modelled according physical properties using data from a range of sources including x-ray diffraction data and NMR. A template molecule is then selected onto which chemical groups which mimic the pharmacophore can be added. The selection can be made such that the mimetic is easy to synthesise, likely to be pharmacologically acceptable, does degrade in vivo and retains the biological activity of the lead compound. Further optimisation or modification can be carried out to select one or more final mimetics useful for in vivo or clinical testing.

It is also possible to isolate a target-specific antibody and then solve its crystal structure. In principle, this approach yields a pharmacophore upon which

subsequent drug design can be based as described above. It may be possible to avoid protein crystallography altogether by generating anti-idiotypic antibodies (antiids) to a functional, pharmacologically active antibody. As a mirror image of a mirror image, the binding site of

- 20 -

the anti-ids would be expected to be an analogue of the original receptor. The anti-id could then be used to isolate peptides from chemically or biologically produced peptide banks.

Any of the therapeutic methods described above may be applied to any subject in need of such therapy, including, for example, mammals such as dogs, cats, cows, horses, rabbits, monkeys, and most preferably, humans.

15 Diagnostic applications

10

20

25

30

35

Polynucleotide sequences of the invention may be used for the diagnosis of epilepsy, in particular generalised epilepsy with febrile seizures plus, as well as other disorders associated with sodium channel dysfunction, including but not restricted to, malignant hyperthermia, myasthenia, episodic ataxia, neuropathic and inflammatory Alzheimer's disease, Parkinson's pain, schizophrenia, hyperekplexia, myotonias such as hypo- and hyperkalaemic periodic paralysis, paramyotonia congenita and potassium aggravated myotonia as well as cardiac arrhythmias such as long QT syndrome, and the use of the DNA molecules of the invention in diagnosis of these disorders, is therefore contemplated.

In another embodiment of the invention, the polynucleotides that may be used for diagnostic purposes include oligonucleotide sequences, genomic DNA and complementary RNA and DNA molecules. The polynucleotides may be used to detect and quantitate gene expression in biological samples. Genomic DNA used for the diagnosis may be obtained from body cells, such as those present in the blood, tissue biopsy, surgical specimen, or autopsy material. The DNA may be isolated and used directly for

- 21 -

detection of a specific sequence or may be amplified by the polymerase chain reaction (PCR) prior to analysis. Similarly, RNA or cDNA may also be used, with or without PCR amplification. To detect a specific nucleic acid sequence, hybridisation using specific oligonucleotides, restriction enzyme digest and mapping, PCR mapping, RNAse protection, and various other methods may be employed. For instance direct nucleotide sequencing of amplification products from the sodium channel subunits can be employed. Sequence of the sample amplicon is compared to that of the wild-type amplicon to determine the presence (or absence) of nucleotide differences.

10

15

20

25

30

35

According to a further aspect of the invention there is provided the use of a polypeptide as described above in the diagnosis of epilepsy, in particular generalised epilepsy with febrile seizures plus, as well as other disorders associated with sodium channel dysfunction, as described above.

When a diagnostic assay is to be based upon mutant proteins constituting a sodium channel, a variety of approaches are possible. For example, diagnosis can be achieved by monitoring differences in the electrophoretic mobility of normal and mutant alpha subunit proteins that form part of the sodium channel. Such an approach will be particularly useful in identifying mutants in which charge in which substitutions are present, or insertions, deletions or substitutions have resulted in a significant change in the electrophoretic migration of the resultant protein. Alternatively, diagnosis may be based upon differences in the proteolytic cleavage patterns of normal and mutant proteins, differences in molar ratios of the various amino acid residues, or by functional assays demonstrating altered function of the gene products.

In another aspect, antibodies that specifically bind mutant sodium channels may be used for the diagnosis of epilepsy, or in assays to monitor patients being treated with agonists, antagonists, modulators or inhibitors of

- 22 -

the mutant sodium channel. Antibodies useful for diagnostic purposes may be prepared in the same manner as described above for therapeutics. Diagnostic assays to detect mutant sodium channels include methods that utilize the antibody and a label to detect a mutant sodium channel in human body fluids or in extracts of cells or tissues. The antibodies may be used with or without modification, and may be labelled by covalent or non-covalent attachment of a reporter molecule.

A variety of protocols for measuring the presence of mutant sodium channels, including ELISAs, RIAs, and FACS, are known in the art and provide a basis for diagnosing epilepsy, in particular generalised epilepsy with febrile seizures plus, as well as other disorders associated with sodium channel dysfunction, as described above. expression of a mutant channel is established by combining body fluids or cell extracts taken from test mammalian subjects, preferably human, with antibody to the channel under conditions suitable for complex formation. amount of complex formation may be quantitated by various methods, preferably by photometric means. Antibodies specific for the mutant channel will only bind to individuals expressing the said mutant channel and not to individuals expressing only wild-type channels (ie normal individuals). This establishes the basis for diagnosing the disease.

Once an individual has been diagnosed with the disorder, effective treatments can be initiated. These may include administering a selective modulator of the mutant channel or an antagonist to the mutant channel such as an antibody or mutant complement as described above. Alternative treatments include the administering of a selective agonist or modulator to the mutant channel so as to restore channel function to a normal level.

35

30

10

15

20

25

Microarray

In further embodiments, complete cDNAs,

- 23 -

oligonucleotides or longer fragments derived from any of the polynucleotide sequences described herein may be used as probes in a microarray. The microarray can be used to monitor the expression level of large numbers of genes identify simultaneously and to genetic variants, mutations, and polymorphisms. This information may be used to determine gene function, to understand the genetic basis of a disorder, to diagnose a disorder, develop and monitor the activities of therapeutic agents. Microarrays may be prepared, used, and analyzed using methods known in the art. (For example, see Schena et al., 1996; Heller et al., 1997).

According to a further aspect of the present neurological material obtained from models generated as a result of the identification of specific sodium channel alpha subunit human mutations, particularly those disclosed in the present invention, can be used in microarray experiments. These experiments can be conducted to identify the level of expression of specific sodium channel alpha subunits, or any cDNA clones from whole-brain libraries, in epileptic brain tissue as opposed to normal control brain tissue. Variations in the expression level of genes, including sodium channel alpha the two tissues indicates between involvement in the epileptic process either as a cause or consequence of the original sodium channel mutation present in the animal model. Microarrays may be prepared, as described above.

30 Transformed hosts

10

15

20

25

35

The present invention also provides for the production of genetically modified (knock-out, knock-in and transgenic), non-human animal models transformed with the DNA molecules of the invention. These animals are useful for the study of the function of a sodium channel, to study the mechanisms of disease as related to a sodium channel, for the screening of candidate pharmaceutical

WO 02/50096

5

10

15

20

25

30

35

- 24 -

PCT/AU01/01648

compounds, for the creation of explanted mammalian cell cultures which express a mutant sodium channel and for the evaluation of potential therapeutic interventions.

Animal species which are suitable for use in the animal models of the present invention include, but are not limited to, rats, mice, hamsters, guinea pigs, rabbits, dogs, cats, goats, sheep, pigs, and non-human primates such as monkeys and chimpanzees. For initial studies, genetically modified mice and rats are highly desirable due to their relative ease of maintenance and shorter life spans. For certain studies, transgenic yeast or invertebrates may be suitable and preferred because they allow for rapid screening and provide for much easier handling. For longer term studies, non-human primates may be desired due to their similarity with humans.

To create an animal model for a mutated sodium channel several methods can be employed. These include but are not limited to generation of a specific mutation in a homologous animal gene, insertion of a wild type human and/or a humanized animal gene by homologous recombination, insertion of a mutant (single or multiple) human gene as genomic or minigene cDNA constructs using wild type or mutant or artificial promoter elements or of artificially modified fragments of insertion the homologous endogenous gene by recombination. The modifications include insertion of mutant stop codons, the deletion of DNA sequences, orthe inclusion of recombination elements (lox p sites) recognized by enzymes such as Cre recombinase.

To create a transgenic or gene targeted (knock-in) mouse, which are preferred, a mutant version of a sodium channel alpha subunit can be inserted into a mouse germ line using standard techniques of occyte microinjection, or transfected into embryonic stem cells, respectively. Alternatively, if it is desired to inactivate or replace an endogenous sodium channel alpha subunit gene, homologous recombination using embryonic stem cells may be

- 25 -

applied.

WO 02/50096

10

15

20

25

30

35

For occyte injection, one or more copies of the mutant sodium channel alpha subunit gene can be inserted into the pronucleus of a just-fertilized mouse occyte. This occyte is then reimplanted into a pseudo-pregnant foster mother. The liveborn mice can then be screened for integrants using analysis of tail DNA or DNA from other tissues for the presence of the particular human subunit gene sequence. The transgene can be either a complete genomic sequence injected as a YAC, BAC, PAC or other chromosome DNA fragment, a complete cDNA with either the natural promoter or a heterologous promoter, or a minigene containing all of the coding region and other elements found to be necessary for optimum expression.

PCT/AU01/01648

According to still another aspect of the invention there is provided the use of genetically modified non-human animals as described above for the screening of candidate pharmaceutical compounds.

It will be clearly understood that, although a number of prior art publications are referred to herein, this reference does not constitute an admission that any of these documents forms part of the common general knowledge in the art, in Australia or in any other country.

Throughout this specification and the claims, the words "comprise", "comprises" and "comprising" are used in a non-exclusive sense, except where the context requires otherwise.

Brief Description of the Drawings

Preferred forms of the invention are described, by way of example only, with reference to the following examples and the accompanying drawings, in which:

Figure 1. Generalised epilepsy with febrile seizures plus (GEFS+) pedigrees are shown for the three families. DNA was not available from those individuals not assigned a letter (X, Y, or Z) or a 0. A: Pedigree of an Australian family with individual numbering for this

- 26 -

family based on Figure 1 in Scheffer & Berkovic (1997). B: Pedigree of an Ashkenazi family. C: Pedigree of a Druze family.

Figure 2. Schematic of the alpha subunit of the sodium channel (SCN1A), showing the position of the three mutations identified in this study.

Figure 3. Sodium channel amino acid alignments. Alignment of sodium channel amino acids surrounding the three SCN1A mutations.

10

15

20

25

30

Modes for Performing the Invention

Example 1: Clinical diagnosis of affected family members

A group of 53 unrelated probands with GEFS+ phenotypes were studied. These subjects were ascertained on the basis of twin and family studies and on the basis of routine clinical practice. Phenotypes in probands and family members were classified as described elsewhere (Scheffer & Berkovic 1997; Singh et al 1999). Familial cases (n=36) were those in which at least one first-degree relative of the proband had a phenotype within the GEFS+ spectrum. Informed consent was obtained from all subjects.

The Australian family in Figure 1A, which has been described extensively elsewhere (Scheffer & Berkovic, 1997; Lopes-Cendes et al, 2000), is the original pedigree leading to the initial delineation and description of the GEFS+ syndrome.

The Israeli family in Figure 1B is of Ashkenazi origin and spans six generations. Twelve family members had seizures. In the two oldest members (I-2, III-3) seizures had occurred in childhood but the data were insufficient to allow classification of the phenotype. Of the 10 other family members who had seizures, 3 had febrile seizures with onset at age 9-13 months. All attacks occurred with fever and offset occurred between 1 and 4 years with 1 to 7 attacks each. Five had febrile seizures plus with onset at age 9-24 months, offset between 5 and 41 years and 2 to 15 attacks each. Seizures

5

10

15

20

25

30

35

- 27 -

during childhood were a mixture of febrile seizures and afebrile tonic-clonic seizures, whereas occurring seizures during teenage and adult years were all Subject V-16 had a more severe phenotype with approximately 20 febrile seizures at age 6 months to 5 years, 10 afebrile tonic-clonic seizures at age 5 to 15 years and occasional complex partial seizures associated with mild learning difficulties. She was classified as having febrile seizures plus and complex partial seizures. Her older sister (V-15) had typical febrile seizures plus, but their younger brother (V-17), aged 14 years, had no febrile seizures but had two afebrile tonic-clonic seizures at ages 12 years 6 months and 14 years. For purposes of linkage analysis, he was regarded as affected, although he had only afebrile tonic-clonic seizures. All affected subjects were of normal or superior intellect, except V-16 (see above) and all had a normal neurological examination. Electroencephalography (EEG) studies had been performed infrequently during the active phase of the epilepsy, and the results usually either were normal or were reported to show generalised discharges.

The second Israeli family was of Druze origin; the parents were from different but proximate villages and were not known to be related. This family spans two generations, and four family members had seizures (Figure 1C). The proband aged 41 years (I-2) had had hundreds of tonic-clonic seizures, sometimes with fever. These began at age 4 years and continued, at a rate of approximately one per month, until the time of the study. The proband was mildly intellectually impaired. EEG showed generalized irregular spike-wave and polyspike-wave discharges, and febrile seizures plus was diagnosed. Of her four children, the oldest was unaffected (II-1), two had febrile seizures (II-2, II-4) and one had febrile seizures plus (II-3).

Example 2: Isolation and sequencing of SCN1A genomic clones

5

10

15

20

25

30

35

- 28 -

At the commencement of this study the full-length sequence of the human SCN1A gene was not known. To determine this sequence a human BAC library obtained from Genome Systems was initially screened to identify human genomic sequence clones containing the SCN1A gene. The BAC filters were screened with a PCR product amplified with the primer pair 5' AGATGACCAGAGTGAATATGTGACTAC 3' (SEQ ID NO:13) and 5' CCAATGGTAAAATAATAATGGCGT 3' (SEQ ID NO:14) designed from the partial cDNA sequence of human SCN1A (Genbank Accession Number X65362).

The BAC filters were hybridised and washed according to manufacturers recommendations. Initially, membranes were individually pre-hybridised in large glass bottles for at least 2 hours in 20 ml of 6X SSC; 0.5% SDS; 5X Denhardt's; 100 ug/ml denatured salmon sperm DNA at 65°C. Overnight hybridisations with $[\alpha^{-32}P]dCTP$ labelled probes were performed at 65°C in 20 ml of a solution containing 6X SSC; 0.5% SDS; 100 ug/ml denatured salmon sperm DNA. Filters were washed sequentially in solutions of 2X SSC; 0.5% SDS (room temperature 5 minutes), 2X SSC; 0.1% SDS (room temperature 15 minutes) and 0.1X SSC; 0.5% SDS (37°C 1 hour if needed).

A number of BAC clones were identified from this hybridisation and BAC129e04 was selected for subcloning and sequencing. DNA from this BAC clone was sheared by nebulisation (10psi for 45 seconds). Sheared DNA was then blunt ended using standard methodologies (Sambrook et al., 1989) and run on an agarose gel in order to isolate DNA in the 2-4 Kb size range. These fragments were cleaned from the agarose using QIAquick columns (Qiagen), ligated into puc18 and used to transform competent XL-1 Blue E. colicells. DNA was isolated from transformed clones and was sequenced using vector specific primers on an ABI377 sequencer to generate 1X coverage of the BAC clone. Sequence data were assembled in contigs using the Phred, Phrap and Gap4 high throughput sequencing software. Exonintron boundaries were predicted based on the rat Scn1a

- 29 -

cDNA sequence (Genbank Accession Number M22253) due to the full length human cDNA sequence of SCN1A not being known.

The human SCN1A gene was determined to be 8,381 base pair in length and is organised into 27 exons spanning over 100 Kb of genomic DNA. To facilitate a comparison with related sodium channels SCN4A, SCN5A and SCN8A, the first untranslated exon of SCN1A is designated exon 1A and the second exon, containing the start codon, remains exon 1 (Table 1). The SCN1A gene shows high homology to SCN2A and SCN3A at both the DNA and protein level. The close proximity of these genes to each other on chromosome 2 indicates likely duplication events during the evolution of the sodium channel gene family. Compared to SCN4A and SCN8A, additional sequence is present in the 3'UTR of SCN1A, giving the final exon an overall length of ~3.3 Kb.

10

15

20

25

30

35

Inspection of the splice junctions of SCN1A shows that there is close agreement with consensus splice motifs, with all introns bounded by GT-AG, except for two (introns 2 and 23). These introns exhibit deviation from the consensus splice pattern and are bounded by AT-AC terminal dinucleotides. These rare splice site variations are conserved in other characterised sodium channel subunits (SCN4A, SCN8A and the more distantly related SCN5A), indicating their ancient origin.

The intron positions are also highly conserved between sodium channel subunits, with most variation seen in the region that codes for the cytoplasmic loop between domains I and II of the gene (Table 1). Within this region, alternative splicing of exon 11 of SCN1A was found that was comparable to the alternative splicing of exon 10B in SCN8A (Plummer et al. 1998). Cytoplasmic loop 1 varies in both length and composition and is the proposed site of functional diversity among different sodium channels (Plummer & Meisler, 1999).

Example 3: Analysis of SCN1A for mutations in epilepsy

The determination of the genomic structure of SCN1A

10

15

20

25

30

35

- 30 -

allowed the design of intronic primers (Table 2 and SEQ ID Numbers:15-88) to amplify each of the 27 exons of SCN1A in order to test for mutations in patients with generalised epilepsy with febrile seizures plus (GEFS+). A total of 53 unrelated patients (as described above) were screened by fluorescent single stranded conformation polymorphism (SSCP) analysis.

HEX-labelled primers were designed to amplify all exons of SCN1A (Table 2). A 30 ng sample of patient DNA was amplified in a total volume of 10 ul. Products were non-denaturing polyacrylamide separated on 4% containing 2% glycerol using the GelScan 2000 (Corbett Research). PCR products showing a conformational change reamplified from 100 ng of genomic unlabelled primers andsequenced using the BigDye Terminator ready reaction kit (Perkin Elmer) according to manufacturers instructions.

A total of 53 unrelated patients with GEFS+ were screened by fluorescent SSCP, including two families consistent with mapping to the same location as SCN1A on chromosome 2 (Figures 1A and 1B). No mutations were found in 17 sporadic cases of GEFS+ that were tested. Of the 36 families tested, 3 were found to have point mutations in SCN1A, which alter the amino acid sequence and are not present in the control population (n=60). The phenotype in the family in Figure 1A previously had been mapped to chromosome 2 (Lopes-Cendes et al. 2000) and carries an A to T mutation at position 563 of the SCN1A coding sequence. This mutation segregates with affected family members. This mutation in exon 4 of SCN1A results in a D188V amino acid substitution that lies just outside the S3 segment of domain I (Figure 2). The aspartic acid residue is conserved in all identified sodium channels in humans as well as in many different animal species, except the jellyfish which has an arginine at this residue and the flatworm which has a serine (Figure 3). The published rat Scn2a sequence (Genbank Accession Number NM_012647)

- 31 -

also has an arginine in place of the aspartic acid at residue 188.

A mutation in exon 21 (G to C nucleotide change at position 4057 of the SCN1A coding sequence) was found to segregate with GEFS+ in the Ashkenazi family (Figure 1B). This mutation changes a highly conserved amino acid (V1353L) located in the S5 segment of domain III (Figure 2). One family member (V-13) did not carry the mutation (Figure 1B). This was determined by testing the DNA of a parent of this family member, since the subjects DNA was unavailable. This individual, who had typical febrile seizures that terminated at an early age, is likely to be a phenocopy. Mutations in the S5 segment of SCN4A that cause hyperkalemic periodic paralysis have been shown also to affect the rate of channel inactivation (Bendahhou et al., 1999)

10

15

20

25

30

A third mutation (C to G nucleotide change at position 4968 of the SCN1A coding sequence) discovered in the Druze family (Figure 1C), changes an amino acid (I1656M) in the S4 segment of domain IV (Figure 2). The S4 segment has a role in channel gating and mutations in this region of SCN1A reduce the rate of inactivation (Kuhn and Greef, 1996).

During the mutation screen of SCN1A several single nucleotide polymorphisms (SNPs) were identified (Table 3). The R1928G variant was found at low frequency in both GEFS+ and control populations. The T1067A variant was common in both populations and the remaining SNPs identified did not alter the amino acid sequence of SCN1A (Table 3).

Example 4: Analysis of a mutated sodium channels and sodium channel alpha subunits

The following methods are used to determine the structure and function of mutated sodium channel or sodium channel alpha subunits.

- 32 -

PCT/AU01/01648

Molecular biological studies

WO 02/50096

10

15

20

25

30

The ability of the mutated sodium channel as a whole or through individual alpha subunits to bind known and unknown proteins can be examined. Procedures such as the yeast two-hybrid system are used to discover and identify any functional partners. The principle behind the yeast procedure is that many eukaryotic two-hybrid transcriptional activators, including those in yeast, consist of two discrete modular domains. The first is a DNA-binding domain that binds to a specific promoter sequence and the second is an activation domain that directs the RNA polymerase II complex to transcribe the gene downstream of the DNA binding site. Both domains are required for transcriptional activation as neither domain can activate transcription on its own. In the yeast twohybrid procedure, the gene of interest or parts thereof (BAIT), is cloned in such a way that it is expressed as a fusion to a peptide that has a DNA binding domain. A second gene, or number of genes, such as those from a cDNA library (TARGET), is cloned so that it is expressed as a fusion to an activation domain. Interaction of the protein of interest with its binding partner brings the DNAbinding peptide together with the activation domain and initiates transcription of the reporter genes. The first reporter gene will select for yeast cells that contain proteins (this reporter is usually interacting nutritional gene required for growth on selective media). The second reporter is used for confirmation and while being expressed in response to interacting proteins it is usually not required for growth.

The nature of the genes and proteins interacting with the mutant sodium channels can also be studied such that these partners can also be targets for drug discovery.

35 Structural studies

Recombinant proteins corresponding to mutated sodium channel alpha subunits can be produced in bacterial,

- 33 -

yeast, insect and/or mammalian cells and used in crystallographical and NMR studies. Together with molecular modeling of the protein, structure-driven drug design can be facilitated.

5

10

15

20

25

30

35

Example 5: Generation of polyclonal antibodies against a mutant sodium channel or sodium channel alpha subunit

Following the identification of new mutations in the alpha subunit of the sodium channel in individuals with generalised epilepsy with febrile seizures plus, antibodies can be made to the mutant channel which can selectively bind and distinguish mutant from normal protein. Antibodies specific for mutagenised epitopes are especially useful in cell culture assays to screen for cells which have been treated with pharmaceutical agents to evaluate the therapeutic potential of the agent.

To prepare polyclonal antibodies, short peptides can be designed homologous to a sodium channel subunit amino acid sequence. Such peptides are typically 10 to 15 amino acids in length. These peptides should be designed in regions of least homology to other receptor subunits and should also have poor homology to the mouse orthologue to avoid cross species interactions in further down-stream experiments such as monoclonal antibody production. Synthetic peptides can then be conjugated to biotin (Sulfo-NHS-LC Biotin) using standard protocols supplied with commercially available kits such as the PIERCETM kit (PIERCE). Biotinylated peptides are subsequently complexed with avidin in solution and for each peptide complex, 2 rabbits are immunized with 4 doses of antigen (200 ug per dose) in intervals of three weeks between doses. initial dose is mixed with Freund's Complete adjuvant while subsequent doses are combined with Freund's Immunoadjuvant. After completion of the immunization, rabbits are test bled and reactivity of sera is assayed by dot blot with serial dilutions of the original peptides. If rabbits show significant reactivity compared with pre-

- 34 -

immune sera, they are then sacrificed and the blood collected such that immune sera can be separated for further experiments.

This procedure is repeated to generate antibodies of wild-type forms receptor subunits. The against antibodies specific for mutant sodium channels can subsequently be used to detect the presence and the relative level of the mutant forms in various tissues.

Example 6: Generation of monoclonal antibodies against a mutant sodium channel or sodium channel alpha subunit

antibodies can Monoclonal be prepared in the following manner. Immunogen, comprising intact mutated sodium channel or sodium channel alpha subunit peptides, is injected in Freund's adjuvant into mice with each mouse receiving four injections of 10 ug to 100 ug of immunogen. After the fourth injection blood samples taken from the mice are examined for the presence of antibody to the Immune mice are sacrificed, immunogen. their removed and single cell suspensions are prepared (Harlow and Lane, 1988). The spleen cells serve as a source of lymphocytes, which are then fused with a permanently growing myeloma partner cell (Kohler and Milstein, 1975). Cells are plated at a density of 2X105 cells/well in 96 well plates and individual wells are examined for growth. These wells are then tested for the presence of sodium channel specific antibodies by ELISA or RIA using wild type or mutant subunit target protein. Cells in positive wells are expanded and subcloned to establish and confirm monoclonality. Clones with the desired specificity are expanded and grown as ascites in mice followed by purification using affinity chromatography using Protein A Sepharose, ion-exchange chromatography or variations and combinations of these techniques.

Industrial Applicability

5

10

15

20

25

30

35

The present invention allows for the diagnosis and

- 35 -

treatment of epilepsy or other disorders associated with sodium channel dysfunction, including but not restricted to, malignant hyperthermia, myasthenia, episodic ataxia, neuropathic and inflammatory pain, Alzheimer's disease, disease, schizophrenia, hyperekplexia, 5 Parkinson's myotonias such as hypo- and hyperkalaemic periodic paralysis, paramyotonia congenita and potassium aggravated myotonia as well as cardiac arrhythmias such as long QT syndrome. In particular, the present invention allows for the diagnosis and treatment of generalised epilepsy with febrile seizures plus.

TABLE 1

- 36 -

Comparison of Exon Sizes of SCN1A with Other Human SCNA Subunits

SCI	N1A	SC	N4A	SC	CN8A	SC	N5A
Exon	Exon	Exon	Exon	Exon	Exon	Exon	Exon
No.	Size	No.	Size	No.	Size	No.	Size
1A	217	-	_	_	_	1	98
1	313	1	661	1	2 7 6	2	324
2	19	2	119	2	121	3	119
3	90	3	90	3	88	4	90
4	20	A	129	4	129	5	129
5	DI		92	5	92	6	92
6			333	6	222	7	231
7			64	7	64	8	64
8			142	8	142	9	142
9	T		210	9	207	10	198
10			154	10A	294	11	180
11	C loc	pl	-	10B	396	12	372
12	L		_	10C	133	13	133
13			239	11	239	14	239
14	DII		174	12	174	15	174
15			357	13	357	16	351
16		•	477	14	471	17	441
	C 100	p2				18	162
17	Ĺ		136	15	118	19	121
18			155	16	155	20	155
19		_	174	17	174	21	174
20	D III	Ī.	123	18A	123	22	123
21			279	19	285	23	282
22	T		54	20	54	24	54
23			138	21	138	25	138
24			105	22	105	26	105
25	D IV		271	23	271	27	271
26	4	<u>.</u>	>2242	24	>1158	28	3257

Note: D: Transmembrane domain; C: Cytoplasmic loop.

- 37 -

Primer Sequences Used for Mutation Analysis of SCN1A

TABLE 2

Exon	Forward Primer	Reverse Primer	Size (bp)
1A	TACCATAGAGTGAGGCGAGG	ATGGACTTCCTGCTCTGCCC	356
1	CCTCTAGCTCATGTTTCATGAC	TGCAGTAGGCAATTAGCAGC	448
2	CTAATTAAGAAGAGATCCAGTGACAG	GCTATAAAGTGCTTACAGATCATGTAC	356
3	CCCTGAATTTTGGCTAAGCTGCAG	CTACATTAAGACACAGTTTCAAAATCC	263
4	GGGCTACGTTTCATTTGTATG	GCAACCTATTCTTAAAGCATAAGACTG	355
5	AGGCTCTTTGTACCTACAGC	CATGTAGGGTCCGTCTCATT	199
6	CACACGTGTTAAGTCTTCATAGT	AGCCCCTCAAGTATTTATCCT	394
7	GAACCTGACCTTCCTGTTCTC	GTTGGCTGTTATCTTCAGTTTC	241
8	GACTAGGCAATATCATAGCATAG	CTTTCTACTATATTATCATCCGG	320
9	TTGAAAGTTGAAGCCACCAC	CCACCTGCTCTTAGGTACTC	363
10	GCCATGCAAATACTTCAGCCC	CACAACAGTGGTTGATTCAGTTG	480
11a	TGAATGCTGAAATCTCCTTCTAC	CTCAGGTTGCTGTTGCGTCTC	306
11b	GATAACGAGAGCCGTAGAGAT	TCTGTAGAAACACTGGCTGG	315
12	CATGAAATTCACTGTGTCACC	CAGCTCTTGAATTAGACTGTC	347
13a	ATCCTTGGGAGGTTTAGAGT	CATCACAACCAGGTTGACAAC	292
13b	CTGGGACTGTTCTCCATATTG	GCATGAAGGATGGTTGAAAG	277
14	CATTGTGGGAAAATAGCATAAGC	GCTATGCAGAACCCTGATTG	338
15a	TGAGACGGTTAGGGCAGATC	AGAAGTCATTCATGTGCCAGC	348
15b	CTGCAAGATCGCCAGTGATTG	ACATGTGCACAATGTGCAGG	276
16a	GTGGTGTTTCCTTCTCATCAAG	TCTGCTGTATGATTGGACATAC	387
16b	CAACAGTCCTTCATTAGGAAAC	ACCTTCCCACACCTATAGAATC	3 5 3
17	CTTGGCAGGCAACTTATTACC	CAAGCTGCACTCCAAATGAAAG	232
18	TGGAAGCAGAGACACTTTATCTAC	GTGCTGTATCACCTTTTCTTAATC	234
19	CCTATTCCAATGAAATGTCATATG	CAAGCTACCTTGAACAGAGAC	318
20	CTACACATTGAATGATGATTCTGT	GCTATATACAATACTTCAGGTTCT	216
21a	ACCAGAGATTACTAGGGGAAT	CCATCGAGCAGTCTCATTTCT	303
21b	ACAACTGGTGACAGGTTTGAC	CTGGGCTCATAAACTTGTACTAAC	297
22	ACTGTCTTGGTCCAAAATCTG	TTCGATTAATTTTACCACCTGATC	267
23	AGCACCAGTGACATTTCCAAC	GGCAGAGAAAACACTCCAAGG	272
24	GACACAGTTTTAACCAGTTTG	TGTGAGACAAGCATGCAAGTT	207
25	CAGGGCCAATGACTACTTTGC	CTGATTGCTGGGATGATCTTGAATC	477
26a	CGCATGATTTCTTCACTGGTTGG	GCGTAGATGAACATGACTAGG	247
26b	TCCTGCGTTGTTTAACATCGG	ATTCCAACAGATGGGTTCCCA	288
26c	TGGAAGCTCAGTTAAGGGAGA	AGCGCAGCTGCAAACTGAGAT	261
26d	CCGATGCAACTCAGTTCATGGA	GTAGTGATTGGCTGATAGGAG	274
26e	AGAGCGATTCATGGCTTCCAATCC	$\mathtt{TGCCTTCTTGCTCATGTTTTTCCACA}$	335
26f	CCTATGACCGGGTGACAAAGCC	TGCTGACAAGGGGTCACTGTCT	242

²⁶f CCTATGACCGGGTGACAAGCC TGCTGACAAGGGGTCACTGTCT 242

Note: Primer sequences are listed 5' to 3'. Due to the large size of 5 exons 11, 13, 15, 16, 21 and 26, the exons were split into two or more overlapping amplicons.

5

TABLE 3

SCN1A Polymorphisms Identified									
	SCN1A polymorph	Freque	ncy (%)						
Position	Mutation	Amino Acid Change	GEFS+	Normal					
Intron 13	IVS13-37C>A	_	2.4	8.6					
Exon 14	c.2522C>G	_	2.4	8.6					
Inron 15	IVS15+54A>G	-	36.3	23.6					
Exon 15	c.2889T>C	-	1.2	0.0					
Exon 16	c.3199G>A	T1067A	29.5	30.8					
Exon 26	c.5782C>G	R1928G	1.2	1.7					

Note: Total GEFS+ samples = 53; Total normal samples=60.

References

15

References cited herein are listed on the following pages, and are incorporated herein by this reference.

- Baulac S. et al. (1999). Am. J. Hum. Genet. 65: 1078-1085.
 Bendahhou S. et al. (1999). J. Neurosci. 19: 4762-4771.
 Cole, SP. et al. (1984). Mol. Cell Biol. 62: 109-120.
 Cote, RJ. et al. (1983). Proc. Natl. Acad. Sci. USA 80: 2026-2030.
- 10 Escayg A. et al. (2000). Nature Genet. 24: 343-345.
 Goldman, CK. et al. (1997). Nature Biotechnology 15: 462-466.
 - Harlow, E. and Lane, D. (1988). Antibodies: A Laboratory
 Manual (Cold Spring Harbor Laboratory, Cold Spring
 Harbor, NY).
 - Heller, RA. et al. (1997). Proc. Natl. Acad. Sci. USA 94: 2150-2155.
 - Huse, WD. et al. (1989). Science 246: 1275-1281.
 - Kohler, G. and Milstein, C. (1975). Nature 256: 495-497.
- 20 Kozbor, D. et al. (1985). J. Immunol. Methods 81:31-42.

 Kuhn, FJP. and Greeff, NG. (1996). J. Gen. Physiol. 114:

 167-183.
 - Lopes-Cendes I. et al. (2000). Am. J. Hum. Genet. 66: 698-
- 25 Moulard B. et al. (1999). Am. J. Hum. Genet. 65: 1396-1400.
 - Orlandi, R. et al. (1989). Proc. Natl. Acad. Sci. USA 86: 3833-3837.
 - Peiffer A. et al. (1999). Ann. Neurol. 46: 671-678.
- 30 Plummer NW. et al. (1998). Genomics 54: 287-296.
 - Plummer NW. and Meisler MH. (1999). Genomics 57: 323-331.
 - Sambrook, J. et al. (1989). Molecular cloning: a laboratory manual. Second Edition. (Cold Spring Harbour Laboratory Press, New York).
- Scharf, D. et al. (1994). Results Probl. Cell Differ. 20: 125-162.
 - Scheffer IE. and Berkovic SF. (1997). Brain 120: 479-490.

- 40 -

Scheffer IE. et al. (2000). Ann. Neurol. 47: 840-841.
Schena, M. et al. (1996). Proc. Natl. Acad. Sci. USA 93: 10614-10619.

Singh R. et al. (1999). Ann Neurol. 45: 75-81.

5 Wallace RH. et al. (1998). Nature Genet. 19: 366-370.

Winter, G. et al. (1991). Nature 349: 293-299.

- 41 -

Claims

WO 02/50096

1. An isolated nucleic acid molecule encoding a mutant alpha subunit of a mammalian voltage-gated sodium channel, wherein a mutation event selected from the group consisting of point mutations, deletions, insertions and rearrangements has occurred and said mutation event disrupts the functioning of an assembled sodium channel so as to produce an epilepsy phenotype, with the proviso that the mutation event is not a C2624T transition or a G4943A transition in an alpha-1 subunit.

PCT/AU01/01648

2. An isolated nucleic acid molecule as claimed in claim 1 wherein said mutation event occurs in the nucleotides encoding an intracellular loop.

15

10

3. An isolated nucleic acid molecule as claimed in claim 2 wherein said mutation event occurs in the nucleotides encoding the intracellular loop between transmembrane segments 2 and 3 of domain I.

- 4. An isolated nucleic acid molecule as claimed in claim 3 wherein said mutation event is a point mutation.
- 5. An isolated nucleic acid molecule as claimed in claim
 25 4 wherein said mutation event results in replacement of an
 aspartic acid residue at amino acid position 188 of the
 alpha-1 subunit of a sodium channel.
- 6. An isolated nucleic acid molecule as claimed in claim
 30 5 wherein the aspartic acid residue at amino acid position
 188 of the alpha-1 subunit of a sodium channel is replaced
 by a valine.
- 7. An isolated nucleic acid molecule as claimed in claim
 6 wherein said mutation event is an A to T nucleotide
 substitution at position 563 of the coding sequence of the
 alpha-1 subunit of a sodium channel.

8. An isolated nucleic acid molecule as claimed in claim 7 comprising the nucleotide sequence set forth in SEQ ID NO:1.

- 42 -

- 9. An isolated nucleic acid molecule as claimed in claim 1 wherein said mutation event takes place in the nucleotides encoding an S5 segment of a transmembrane domain.
- 10 10. An isolated nucleic acid molecule as claimed in claim 9 wherein said mutation event occurs in the nucleotides encoding the S5 segment of domain III.
- 11. An isolated nucleic acid molecule as claimed in claim
 15 10 wherein said mutation event is a point mutation.
 - 12. An isolated nucleic acid molecule as claimed in claim 11 wherein said mutation event results in replacement of a valine residue at amino acid position 1353 of the alpha-1 subunit of a sodium channel.
- 13. An isolated nucleic acid molecule as claimed in claim
 12 wherein the valine residue at amino acid position 1353
 of the alpha-1 subunit of a sodium channel is replaced by
 25 a leucine.

20

30

- 14. An isolated nucleic acid molecule as claimed in claim 13 wherein said mutation event is a G to C nucleotide substitution at position 4057 of the coding sequence of the alpha-1 subunit of a sodium channel.
- 15. An isolated nucleic acid molecule as claimed in claim 14 comprising the nucleotide sequence set forth in SEQ ID NO:3.
- 16. An isolated nucleic acid molecule as claimed in claim 1 wherein said mutation event occurs in the nucleotides

- 43 -

encoding an S4 segment of a transmembrane domain.

WO 02/50096

17. An isolated nucleic acid molecule as claimed in claim 16 wherein said mutation event occurs in the nucleotides encoding the S4 segment of domain IV.

PCT/AU01/01648

- 18. An isolated nucleic acid molecule as claimed in claim 17 wherein said mutation event is a point mutation.
- 10 19. An isolated nucleic acid molecule as claimed in claim
 18 wherein said mutation event results in replacement of
 an isoleucine residue at amino acid position 1656 of the
 alpha-1 subunit of a sodium channel.
- 15 20. An isolated nucleic acid molecule as claimed in claim 19 wherein the isoleucine residue at amino acid position 1656 of the alpha-1 subunit of a sodium channel is replaced by a methionine.
- 20 21. An isolated nucleic acid molecule as claimed in claim 20 wherein said mutation event is a C to G nucleotide substitution at position 4968 of the coding sequence of the alpha-1 subunit of a sodium channel.
- 25 22. An isolated nucleic acid molecule as claimed in claim 21 comprising the nucleotide sequence set forth in SEQ ID NO:5.
- 23. An isolated nucleic acid molecule as claimed in any one of claims 1 to 22 in which one or more additional mutation events selected from the group consisting of point mutations, deletions, insertions and rearrangements have occurred.
- 24. An isolated nucleic acid molecule as claimed in claim 23 wherein said one or more additional mutation events are point mutations which result in conservative amino acid

PCT/AU01/01648

substitutions.

WO 02/50096

10

30

- 25. An isolated nucleic acid molecule encoding a mutant alpha subunit of a mammalian voltage-gated sodium channel, event selected from wherein a mutation the consisting of point mutations, deletions, insertions and rearrangements has occurred in an intracellular loop, in the S4 segment of domain IV at nucleotide position 4968 of alpha-1 subunit coding sequence ornucleotide position in the coding sequence of other alpha subunits, or in an S5 segment of a transmembrane domain so as to produce an epilepsy phenotype.
- 26. An isolated nucleic acid molecule consisting of the nucleotide sequence set forth in SEQ ID NO:1.
 - 27. An isolated nucleic acid molecule consisting of the nucleotide sequence set forth in SEQ ID NO:3.
- 20 28. An isolated nucleic acid molecule consisting of the nucleotide sequence set forth in SEQ ID NO:5.
- 29. An isolated nucleic acid molecule selected from the group consisting of DNA molecules comprising the nucleotide sequence set forth in any one of SEQ ID NO:7, 8, 9,11 and 89.
 - 30. An isolated polypeptide, said polypeptide being a mutant alpha subunit of a mammalian voltage-gated sodium channel, wherein a mutation event selected from the group consisting of substitutions, deletions, insertions and rearrangements has occurred and said mutation event disrupts the functioning of an assembled sodium channel so as to produce an epilepsy phenotype, with the proviso that the mutation event is not a T875M transition or a R1648H transition in an alpha-1 subunit.

31. An isolated polypeptide as claimed in claim 30 wherein said mutation event occurs in an intracellular

- 45 -

loop.

5 32. An isolated polypeptide as claimed in claim 30 wherein said mutation event occurs in an intracellular loop between transmembrane segments 2 and 3 of domain I.

- 33. An isolated polypeptide as claimed in claim 30 wherein said mutation event is a substitution.
- 34. An isolated polypeptide as claimed in claim 33 wherein the substitution involves replacement of an aspartic acid residue at position 188 of the alpha-1 subunit of a sodium channel.
 - 35. An isolated polypeptide as claimed in claim 34 wherein the aspartic acid residue is replaced with a valine residue.

20

- 36. An isolated polypeptide as claimed in claim 35 comprising the amino acid sequence set forth in SEQ ID NO:2.
- 25 37. An isolated polypeptide as claimed in claim 30 wherein the mutation event occurs in an S5 segment of a transmembrane domain.
- 38. An isolated polypeptide as claimed in claim 37 wherein said mutation event occurs in the S5 segment of domain III.
 - 39. An isolated polypeptide as claimed in claim 38 wherein said mutation event is a substitution.

35

40. An isolated polypeptide as claimed in claim 39 wherein the substitution involves replacement of a valine

. .

WO 02/50096

15

- 46 -

residue at position 1353 of the alpha-1 subunit of a sodium channel.

PCT/AU01/01648

- 41. An isolated polypeptide as claimed in claim 40 wherein the valine residue is replaced with a leucine residue.
- 42. An isolated polypeptide as claimed in claim 41 comprising the amino acid sequence set forth in SEQ ID 10 NO:4.
 - 43. An isolated polypeptide as claimed in claim 30 wherein said mutation event occurs in an S4 segment of a transmembrane domain.

44. An isolated polypeptide as claimed in claim 41 wherein said mutation event occurs in the S4 segment of domain IV.

- 20 45. An isolated polypeptide as claimed in claim 44 wherein an isoleucine residue at position 1656 of the alpha-1 subunit of a sodium channel is replaced.
- 46. An isolated polypeptide as claimed in claim 45 wherein the isoleucine residue is replaced with a methionine residue.
- 47. An isolated polypeptide as claimed in claim 46 comprising the amino acid sequence set forth in SEQ ID 30 NO:6.
 - 48. An isolated polypeptide, said polypeptide being a mutant α -subunit of a mammalian voltage-gated sodium channel, wherein a mutation event selected from the group of substitutions, deletions, insertions and rearrangements has occurred in an intracellular loop, in the S4 segment of domain IV at amino acid position 1656 of the alpha-1

- 47 -

subunit or homologous amino acid position of other alpha subunits, or in an S5 segment of a transmembrane domain.

- 49. An isolated polypeptide having the amino acid sequence set forth in SEQ ID NO:2.
 - 50. An isolated polypeptide having the amino acid sequence set forth in SEQ ID NO:4.
- 10 51. An isolated polypeptide having the amino acid sequence set forth in SEQ ID NO:6.
- 52. An isolated polypeptide, said polypeptide being an assembled mammalian voltage-gated sodium channel comprising an alpha subunit as defined in any one of claims 30 to 51.
- 53. An isolated polypeptide selected from the group consisting of polypeptides with the amino acid sequence set forth in SEO ID NO:10 or SEQ ID NO:12.
 - 54. A cell transformed with an isolated nucleic acid molecule as claimed in any one of claims 1 to 29.
- 25 55. A cell as claimed in claim 54 which is an eukaryotic cell or bacterial cell.
 - 56. A method of preparing a polypeptide comprising the steps of:
- 30 (1) culturing cells as claimed in claim 54 or 55 under conditions effective for polypeptide production; and
 - (2) harvesting the polypeptide.

35

57. A polypeptide prepared by the method of claim 56.

58. An antibody which is immunologically reactive with a mutant polypeptide as defined in any one of claims 30 to

- 48 -

52, but not with a wild-type mammalian voltage-gated sodium channel.

- 59. An antibody as claimed in claim 58 which is selected from the group consisting of a monoclonal antibody, a humanised antibody, a chimaeric antibody or an antibody fragment including a Fab fragment, (Fab')2 fragment, Fv fragment, single chain antibodies and single domain antibodies.
- 10 60. A method of treating disorders associated with sodium channel dysfunction, comprising administering a selective agonist, antagonist or modulator of the sodium channel when it has undergone a mutation event as defined in any one of claims 30 to 48 to a patient in need of such treatment.
 - 61. The use of a selective agonist, antagonist or modulator of the sodium channel when it has undergone a mutation event as defined in any one of claims 30 to 48 in the manufacture of a medicament for the treatment of a disorder associated with sodium channel dysfunction.

20

25

30

35

62. A method of treating disorders associated with sodium channel dysfunction, comprising administering an isolated DNA molecule which is the complement (antisense) of a nucleic acid molecule as defined in any one of claims 1 to 29 and which encodes an RNA molecule that hybridizes with the mRNA encoding a mutant sodium channel alpha subunit, to a subject in need of such treatment.

63. The use of an isolated DNA molecule which is the complement of a nucleic acid molecule as defined in any one of claims 1 to 29 and which encodes an RNA molecule that hybridizes with the mRNA encoding a mutant sodium channel alpha subunit, in the manufacture of a medicament for the treatment of disorders associated with sodium channel dysfunction.

64. A method of treating disorders associated with sodium channel dysfunction comprising administration of an antibody as defined in claim 58 or 59.

5

15

- 65. Use of a polypeptide as claimed in any one of claims 30 to 53 or 57 for the screening of candidate pharmaceutical agents.
- 10 66. Use as claimed in claim 65 wherein high throughput screening techniques are employed.
 - 67. A genetically modified non-human animal transformed with an isolated nucleic acid molecule as defined in any one of claims 1 to 29.
 - 68. A genetically modified non-human animal as claimed in claim 67 in which the animal is selected from the group consisting of rats, mice, hamsters, guinea pigs, rabbits, dogs, cats, goats, sheep, pigs and non-human primates such as monkeys and chimpanzees.
- 69. The use of a genetically modified non-human animal as claimed in claim 67 or 68 in the screening of candidate pharmaceutical compounds.
 - 70. The use of a cell as claimed in claim 54 to 55 in the screening of candidate pharmaceuticals.
- 30 71. An expression vector comprising a DNA molecule as claimed in any one of claims 1 to 29.
 - 72. A microarray comprising a complete cDNA, an oligonucleotide or a longer fragment derived from any of the polynucleotide sequences defined in claims 1 to 29.
 - 73. The use of a DNA molecule as claimed in any one of

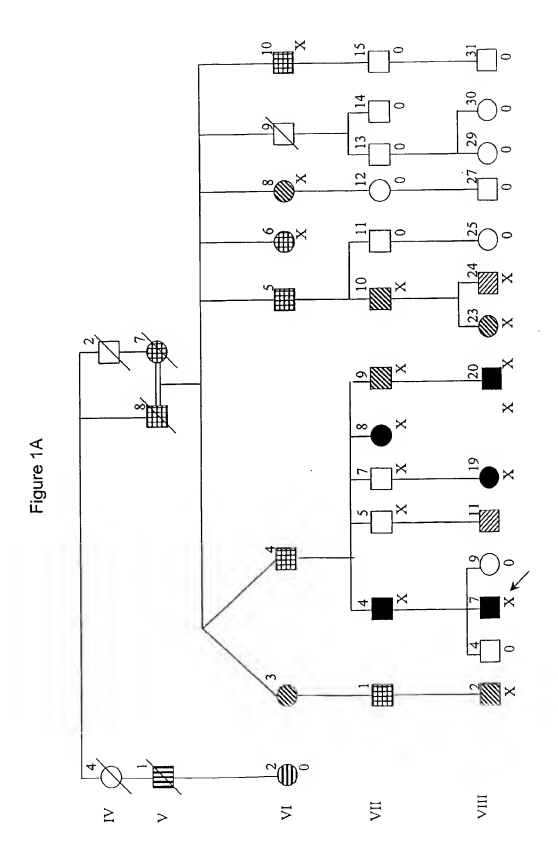
- 50 **-**

claims 1 to 29 in the diagnosis of epilepsy, in particular generalised epilepsy with febrile seizures plus, and other disorders associated with sodium channel dysfunction.

74. The use of a polypeptide as defined in any one of claims 30 to 53 or 57 in the diagnosis of epilepsy, in particular generalised epilepsy with febrile seizures plus, as well as other disorders associated with sodium channel dysfunction.

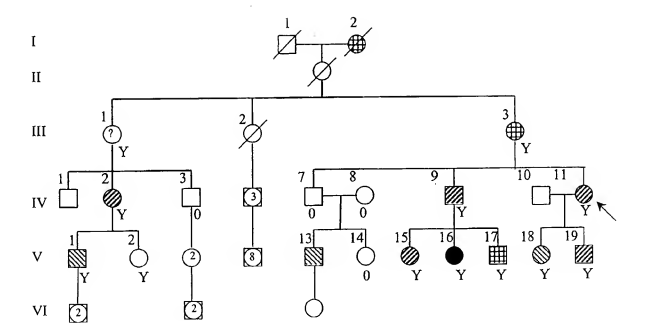
10

75. The use of an antibody as defined in claims 58 or 59 in the diagnosis of epilepsy, in particular generalised epilepsy with febrile seizures plus, as well as other disorders associated with sodium channel dysfunction.



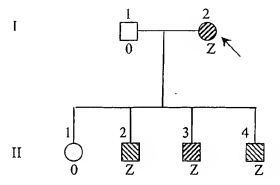
2/5

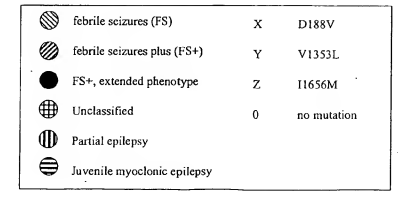
Figure 1B



3/5

Figure 1C





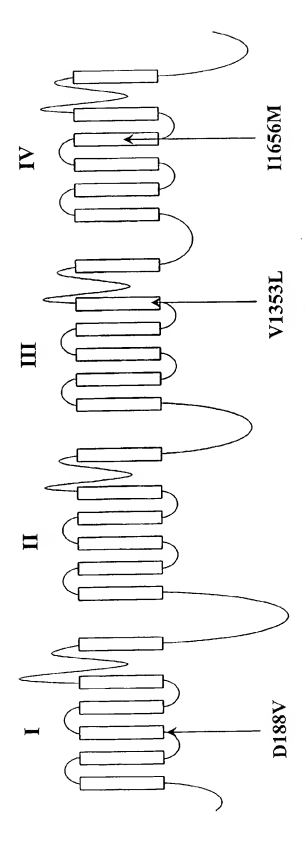


Figure 2

5/5

	د / د	Figure 3
i) D188V SCN1A RAT SCN1A SCN2A SCN3A SCN4A SCN5A SCN6A SCN8A SCN9A SCN10A SCN11A SCN12A EL. EEL DROS SQUID FLATWORM JELLYFISH	F T F L R D	P W N W L
ii) V1353L SCN1A RAT SCN1A SCN2A SCN3A SCN4A SCN5A SCN6A SCN8A SCN9A SCN10A SCN11A SCN12A EL. EEL DROS SQUID FLATWORM JELLYFISH	M N V L L V	C L I F W M I
iii) I1656M SCN1A RAT SCN1A SCN2A SCN3A SCN4A SCN5A SCN6A SCN8A SCN9A SCN10A SCN11A SCN11A SCN12A EL. EEL DROS SQUID FLATWORM JELLYFISH	K G A K G I	

SCN1APCT1.ST25.txt
SEOUENCE LISTING<110> Bionomics Limited<1

20>

P10<130> New SCN1A Mutations<160> 89 <170> PatentIn version 3.1<210> 1<211> 8381<212> DNA<213> Homo sapiens<400> 1 atactgcaga ggtctctggt gcatgtgtgt atgtgtgcgt ttgtgtgtgt ttgtgtgtct

60

gtgtgttctg ccccagtgag actgcagccc ttgtaaatac tttgacacct tttgcaagaa

ggaatctgaa caattgcaac tgaaggcaca ttgttatcat ctcgtctttg ggtgatgctg
180

ttcctcactg cagatggata attttccttt taatcaggaa tttcatatgc agaataaatg 240

gtaattaaaa tgtgcaggat gacaagatgg agcaaacagt gcttgtacca ccaggacctg

acagetteaa ettetteace agagaatete ttgeggetat tgaaagaege attgeagaag 360

aaaaggcaaa gaatcccaaa ccagacaaaa aagatgacga cgaaaatggc ccaaagccaa 420

atagtgactt ggaagctgga aagaaccttc catttattta tggagacatt cctccagaga

tggtgtcaga gccctggag gacctggacc cctactatat caataagaaa acttttatag 540

tattgaataa attgaaggcc atcttccggt tcagtgccac ctctgccctg tacattttaa 600

ctcccttcaa tcctcttagg aaaatagcta ttaagatttt ggtacattca ttattcagca 660

tgctaattat gtgcactatt ttgacaaact gtgtgtttat gacaatgagt aaccctcctg
720

attggacaaa gaatgtagaa tacaccttca caggaatata tacttttgaa tcacttataa 780

SCN1APCT1.ST25.txt

aaattattgc aaggggattc tgtttagaag attttacttt ccttcgggtt ccatggaact 840

ggctcgattt cactgtcatt acatttgcgt acgtcacaga gtttgtggac ctgggcaatg 900

tctcggcatt gagaacattc agagttctcc gagcattgaa gacgatttca gtcattccag

gcctgaaaac cattgtggga gccctgatcc agtctgtgaa gaagctctca gatgtaatga 1020

tcctgactgt gttctgtctg agcgtatttg ctctaattgg gctgcagctg ttcatgggca
1080

acctgaggaa taaatgtata caatggcctc ccaccaatgc ttccttggag gaacatagta 1140

tagaaaagaa tataactgtg aattataatg gtacacttat aaatgaaact gtctttgagt 1200

ttgactggaa gtcatatatt caagattcaa gatatcatta tttcctggag ggttttttag
1260

atgcactact atgtggaaat agctctgatg caggccaatg tccagaggga tatatgtgtg
1320

tgaaagctgg tagaaatccc aattatggct acacaagctt tgataccttc agttgggctt 1380

ttttgtcctt gtttcgacta atgactcagg acttctggga aaatctttat caactgacat 1440

tacgtgctgc tgggaaaacg tacatgatat tttttgtatt ggtcattttc ttgggctcat
1500

tctacctaat aaatttgatc ctggctgtgg tggccatggc ctacgaggaa cagaatcagg
1560

ccaccttgga agaagcagaa cagaaagagg ccgaatttca gcagatgatt gaacagctta

SCN1APCT1.ST25.txt

1620

aaaagcaaca ggaggcagct cagcaggcag caacggcaac tgcctcagaa cattccagag

1680

agcccagtgc agcaggcagg ctctcagaca gctcatctga agcctctaag ttgagttcca

1740

1800

gggaagagaa agatgaggat gaattccaaa aatctgaatc tgaggacagc atcaggagga

1860

aaggttttcg cttctccatt gaagggaacc gattgacata tgaaaagagg tactcctccc

1920

cacaccagtc tttgttgagc atccgtggct ccctattttc accaaggcga aatagcagaa

1980

caagcetttt cagetttaga gggegageaa aggatgtggg atetgagaac gaettegeag

2040

atgatgagca cagcaccttt gaggataacg agagccgtag agattccttg tttgtgcccc

2100

gacgacacgg agagagacgc aacagcaacc tgagtcagac cagtaggtca tcccggatgc

2160

tggcagtgtt tccagcgaat gggaagatgc acagcactgt ggattgcaat ggtgtggttt

2220

ccttggttgg tggaccttca gttcctacat cgcctgttgg acagcttctg ccagaggtga

2280

taatagataa gccagctact gatgacaatg gaacaaccac tgaaactgaa atgagaaaga

2340

gaaggtcaag ttctttccac gtttccatgg actttctaga agatccttcc caaaggcaac

2400

gagcaatgag tatagccagc attctaacaa atacagtaga agaacttgaa gaatccaggc

SCN1APCT1.ST25.txt

2460

agaaatgccc accetgttgg tataaatttt ccaacatatt cttaatctgg gactgttctc 2520

catattggtt aaaagtgaaa catgttgtca acctggttgt gatggaccca tttgttgacc 2580

tggccatcac catctgtatt gtcttaaata ctcttttcat ggccatggag cactatccaa 2640

tgacggacca tttcaataat gtgcttacag taggaaactt ggttttcact gggatcttta
2700

cagcagaaat gtttctgaaa attattgcca tggatcctta ctattatttc caagaaggct 2760

ggaatatett tgacggtttt attgtgacge ttageetggt agaaettgga etegeeaatg 2820

tggaaggatt atctgttctc cgttcatttc gattgctgcg agttttcaag ttggcaaaat 2880

cttggccaac gttaaatatg ctaataaaga tcatcggcaa ttccgtgggg gctctgggaa 2940

atttaaccct cgtcttggcc atcatcgtct tcatttttgc cgtggtcggc atgcagctct 3000

ttggtaaaag ctacaaagat tgtgtctgca agatcgccag tgattgtcaa ctcccacgct 3060

ggcacatgaa tgacttette cacteettee tgattgtgtt eegegtgetg tgtggggagt 3120

ggatagagac catgtgggac tgtatggagg ttgctggtca agccatgtgc cttactgtct 3180

tcatgatggt catggtgatt ggaaacctag tggtcctgaa tctctttctg gccttgcttc 3240

SCN1APCT1.ST25.txt

tgagctcatt tagtgcagac aaccttgcag ccactgatga tgataatgaa atgaataatc 3300

tccaaattgc tgtggatagg atgcacaaag gagtagctta tgtgaaaaga aaaatatatg 3360

aatttattca acagtccttc attaggaaac aaaagatttt agatgaaatt aaaccacttg 3420

atgatctaaa caacaagaaa gacagttgta tgtccaatca tacaacagaa attgggaaag 3480

atottgacta tottaaagat gtaaatggaa ctacaagtgg tataggaact ggcagcagtg 3540

ttgaaaaata cattattgat gaaagtgatt acatgtcatt cataaacaac cccagtctta 3600

ctgtgactgt accaattgct gtaggagaat ctgactttga aaatttaaac acggaagact 3660

ttagtagtga atcggatctg gaagaaagca aagagaaact gaatgaaagc agtagctcat 3720

cagaaggtag cactgtggac atcggcgcac ctgtagaaga acagcccgta gtggaacctg 3780

aagaaactct tgaaccagaa gcttgtttca ctgaaggctg tgtacaaaga ttcaagtgtt 3840

gtcaaatcaa tgtggaagaa ggcagaggaa aacaatggtg gaacctgaga aggacgtgtt 3900

tccgaatagt tgaacataac tggtttgaga ccttcattgt tttcatgatt ctccttagta 3960

gtggtgctct ggcatttgaa gatatata ttgatcagcg aaagacgatt aagacgatgt 4020

tggaatatgc tgacaaggtt ttcacttaca ttttcattct ggaaatgctt ctaaaatggg 4080

Page 5

SCN1APCT1.ST25.txt

tggcatatgg ctatcaaaca tatttcacca atgcctggtg ttggctggac ttcttaattg
4140

ttgatgtttc attggtcagt ttaacagcaa atgccttggg ttactcagaa cttggagcca 4200

tcaaatctct caggacacta agagctctga gacctctaag agccttatct cgatttgaag 4260

ggatgagggt ggttgtgaat gcccttttag gagcaattcc atccatcatg aatgtgcttc 4320

tggtttgtct tatattctgg ctaattttca gcatcatggg cgtaaatttg tttgctggca 4380

aattctacca ctgtattaac accacaactg gtgacaggtt tgacatcgaa gacgtgaata 4440

atcatactga ttgcctaaaa ctaatagaaa gaaatgagac tgctcgatgg aaaaatgtga 4500

aagtaaactt tgataatgta ggatttgggt atctctcttt gcttcaagtt gccacattca 4560

aaggatggat ggatataatg tatgcagcag ttgattccag aaatgtggaa ctccagccta 4620

agtatgaaaa aagtctgtac atgtatcttt actttgttat tttcatcatc tttgggtcct 4680

tcttcacctt gaacctgttt attggtgtca tcatagataa tttcaaccag cagaaaaaga 4740

agtttggagg tcaagacatc tttatgacag aagaacagaa gaaatactat aatgcaatga 4800

aaaaattagg atcgaaaaaa ccgcaaaagc ctatacctcg accaggaaac aaatttcaag
4860

gaatggtctt tgacttcgta accagacaag tttttgacat aagcatCatg attctcatct

SCN1APCT1.ST25.txt

4920

gtcttaacat ggtcacaatg atggtggaaa cagatgacca gagtgaatat gtgactacca 4980

ttttgtcacg catcaatctg gtgttcattg tgctatttac tggagagtgt gtactgaaac 5040

tcatctctct acgccattat tattttacca ttggatggaa tatttttgat tttgtggttg \cdot

tcattctctc cattgtaggt atgtttcttg ccgagctgat agaaaagtat ttcgtgtccc 5160

ctaccctgtt ccgagtgatc cgtcttgcta ggattggccg aatcctacgt ctgatcaaag 5220

gagcaaaggg gatccgcacg ctgctctttg ctttgatgat gtcccttcct gcgttgttta
5280

acateggeet ectaetette etagteatgt teatetaege eatetttggg atgteeaaet 5340

ttgcctatgt taagagggaa gttgggatcg atgacatgtt caactttgag acctttggca 5400

acagcatgat ctgcctattc caaattacaa cctctgctgg ctgggatgga ttgctagcac 5460

ccattctcaa cagtaagcca cccgactgtg accctaataa agttaaccct ggaagctcag 5520

ttaagggaga ctgtgggaac ccatctgttg gaattttctt ttttgtcagt tacatcatca 5580

tatecttect ggttgtggtg aacatgtaca tegeggteat eetggagaac tteagtgttg 5640

ctactgaaga aagtgcagag cctctgagtg aggatgactt tgagatgttc tatgaggttt 5700

gggagaagtt tgatcccgat gcaactcagt tcatggaatt tgaaaaatta tctcagtttg
Page 7

SCN1APCT1.ST25.txt

5760

cagctgcgct tgaaccgcct ctcaatctgc cacaaccaaa caaactccag ctcattgcca

5820

tggatttgcc catggtgagt ggtgaccgga tccactgtct tgatatctta tttgctttta

5880

caaagcgggt tctaggagag agtggagaga tggatgctct acgaatacag atggaagagc

5940

gattcatggc ttccaatcct tccaaggtct cctatcagcc aatcactact actttaaaac

6000

gaaaacaaga ggaagtatet getgteatta tteagegtge ttacagaege cacettttaa

6060

agcgaactgt aaaacaagct tcctttacgt acaataaaaa caaaatcaaa ggtggggcta

6120

atcttcttat aaaagaagac atgataattg acagaataaa tgaaaactct attacagaaa

6180

aaactgatct gaccatgtcc actgcagctt gtccaccttc ctatgaccgg gtgacaaagc

6240

caattgtgga aaaacatgag caagaaggca aagatgaaaa agccaaaggg aaataaatga

6300

aaataaataa aaataattgg gtgacaaatt gtttacagcc tgtgaaggtg atgtattttt

6360

atcaacagga ctcctttagg aggtcaatgc caaactgact gtttttacac aaatctcctt

6420

aaggtcagtg cctacaataa gacagtgacc ccttgtcagc aaactgtgac tctgtgtaaa

6480

ggggagatga ccttgacagg aggttactgt tctcactacc agctgacact gctgaagata

SCN1APCT1.ST25.txt

agatgcacaa tggctagtca gactgtaggg accagtttca aggggtgcaa acctgtgatt 6600

ttggggttgt ttaacatgaa acactttagt gtagtaattg tatccactgt ttgcatttca 6660

actgccacat ttgtcacatt tttatggaat ctgttagtgg attcatcttt ttgttaatcc 6720

atgtgtttat tatatgtgac tatttttgta aacgaagttt ctgttgagaa ataggctaag 6780

gacctctata acaggtatgc cacctggggg gtatggcaac cacatggccc tcccagctac 6840

acaaagtcgt ggtttgcatg agggcatgct gcacttagag atcatgcatg agaaaaagtc 6900

acaagaaaaa caaattotta aatttoacca tatttotggg aggggtaatt gggtgataag 6960

tggaggtgct ttgttgatct tgttttgcga aatccagccc ctagaccaag tagattattt 7020

gtgggtaggc cagtaaatct tagcaggtgc aaacttcatt caaatgtttg gagtcataaa 7080

tgttatgttt ctttttgttg tattaaaaaa aaaacctgaa tagtgaatat tgcccctcac 7140

cctccaccgc cagaagactg aattgaccaa aattactctt tataaatttc tgctttttcc 7200

tgcactttgt ttagccatct ttgggctctc agcaaggttg acactgtata tgttaatgaa 7260

atgctattta ttatgtaaat agtcatttta ccctgtggtg cacgtttgag caaacaaata 7320

atgacctaag cacagtattt attgcatcaa atatgtacca caagaaatgt agagtgcaag 7380

SCN1APCT1.ST25.txt

ctttacacag gtaataaaat gtattctgta ccatttatag atagtttgga tgctatcaat 7440'

gcatgtttat attaccatgc tgctgtatct ggtttctctc actgctcaga atctcattta 7500

tgagaaacca tatgtcagtg gtaaagtcaa ggaaattgtt caacagatct catttattta 7560

agtcattaag caatagtttg cagcacttta acagcttttt ggttattttt acattttaag 7620

tggataacat atggtatata gccagactgt acagacatgt ttaaaaaaac acactgctta 7680

acctattaaa tatgtgttta gaattttata agcaaatata aatactgtaa aaagtcactt 7740

tattttattt ttcagcatta tgtacataaa tatgaagagg aaattatctt caggttgata 7800

tcacaatcac ttttcttact ttctgtccat agtacttttt catgaaagaa atttgctaaa 7860

taagacatga aaacaagact gggtagttgt agatttctgc tttttaaatt acatttgcta .7920

attttagatt atttcacaat tttaaggagc aaaataggtt cacgattcat atccaaatta 7980

tgctttgcaa ttggaaaagg gtttaaaatt ttatttatat ttctggtagt acctgtacta 8040

actgaattga aggtagtgct tatgttattt ttgttctttt tttctgactt cggtttatgt 8100

tttcatttct ttggagtaat gctgctctag attgttctaa atagaatgtg ggcttcataa 8160

tttttttttc cacaaaaaca gagtagtcaa cttatatagt caattacatc aggacatttt

SCN1APCT1.ST25.txt

8220

gtgtttctta cagaagcaaa ccataggctc ctcttttcct taaaactact tagataaact 8280

gtattcgtga actgcatgct ggaaaatgct actattatgc taaataatgc taaccaacat 8340

ttaaaatgtg caaaactaat aaagattaca ttttttattt t

8381

<210> 2<211> 2009<212> PRT<213> Homo sapiens<400> 2

Met Glu Gln Thr Val Leu Val Pro Pro Gly Pro Asp Ser Phe Asn Phe 1 5 10 15

Phe Thr Arg Glu Ser Leu Ala Ala Ile Glu Arg Arg Ile Ala Glu Glu 20 25 30

Lys Ala Lys Asn Pro Lys Pro Asp Lys Lys Asp Asp Asp Glu Asn Gly 35 40 45

Pro Lys Pro Asn Ser Asp Leu Glu Ala Gly Lys Asn Leu Pro Phe Ile 50 55 60

Tyr Gly Asp Ile Pro Pro Glu Met Val Ser Glu Pro Leu Glu Asp Leu 65 70 75 80

Asp Pro Tyr Tyr Ile Asn Lys Lys Thr Phe Ile Val Leu Asn Lys Leu 85 90 95

Lys Ala Ile Phe Arg Phe Ser Ala Thr Ser Ala Leu Tyr Ile Leu Thr 100 105 110

Pro Phe Asn Pro Leu Arg Lys Ile Ala Ile Lys Ile Leu Val His Ser 115 120 125

Leu Phe Ser Met Leu Ile Met Cys Thr Ile Leu Thr Asn Cys Val Phe 130 135 140

Met Thr Met Ser Asn Pro Pro Asp Trp Thr Lys Asn Val Glu Tyr Thr 145 150 155 160

SCN1APCT1.ST25.txt

Phe	Thr	Gly	Ile	Tyr 165	Thr	Phe	Glu	Ser	Leu 170	Ile	Lys	Ile	Ile	Ala 175	Arg
Gly	Phe	Cys	Leu 180	Glu	Asp	Phe	Thr	Phe 185	Leu	Arg	Val	Pro	Trp 190	Asn	Trp
Leu	Asp	Phe 195	Thr	Val	Ile	Thr	Phe 200	Ala	Tyr	Val	Thr	Glu 205	Phe	Val	Asp
Leu	Gly 210	Asn	Val	Ser	Ala	Leu 215	Arg	Thr	Phe	Arg	Val 220	Leu	Arg	Ala	Leu
Lys 225	Thr	Ile	Ser	Val	Ile 230	Pro	Gly	Leu	Lys	Thr 235	Ile	Val	Gly	Ala	Leu 240
Ile	Gln	Ser	Val	Lys 245	Lys	Leu	Ser	Asp	Val 250	Met	Ile	Leu	Thr	Val 255	Phe
Cys	Leu	Ser	Val 260	Phe	Ala	Leu	Ile	Gly 265	Leu	Gln	Leu	Phe	Met 270	Gly	Asn
Leu	Arg	Asn 275	Lys	Cys	Ile	Gln	Trp 280	Pro	Pro	Thr	Asn	Ala 285	Ser	Leu	Glu
Glu	His 290	Ser	Ile	Glu	Lys	Asn 295	Ile	Thr	Val	Asn	Tyr 300	Asn	Gly	Thr	Leu
Ile 305	Asn	Glu	Thr	Val	Phe 310	Glu	Phe	Asp	Trp	Lys 315	Ser	Tyr	Ile	Gln	Asp 320
Ser	Arg	Tyr	His	Tyr 325	Phe	Leu	Glu	Gly	Phe 330	Leu	Asp	Ala	Leu	Leu 335	Cys
Gly	Asn	Ser	Ser 340	Asp	Ala	Gly	Gln	Cys 345	Pro	Glu	Gly	Tyr	Met 350	Cys	Val
Lys	Ala	Gly 355	Arg	Asn	Pro	Asn	Tyr 360	Gly	Tyr	Thr	Ser	Phe 365	Asp	Thr	Phe
Ser	Trp	Ala	Phe	Leu	Ser	Leu	Phe P	Arg age		Met	Thr	Gln	Asp	Phe	Trp

	SCN1APCT1.ST25.txt
370	375

Glu 385	Asn	Leu	Tyr	Gln	Leu 390	Thr	Leu	Arg	Ala	Ala 395	Gly	Lys	Thr	Tyr	Met 400
Ile	Phe	Phe	Val	Leu 405	Val	Ile	Phe	Leu	Gly 410	Ser	Phe	Tyr	Leu	Ile 415	Asn
Leu	Ile	Leu	Ala 420	Val	Val	Ala	Met	Ala 425	Tyr	Glu	Glu	Gln	Asn 430	Gln	Ala
Thr	Leu	Glu 435	Glu	Ala	Glu	Gln	Lys 440	Glu	Ala	Glu	Phe	Gln 445	Gln	Met	Ile
Glu	Gln 450	Leu	Lys	Lys	Gln	Gln 455	Glu	Ala	Ala	Gln	Gln 460	Ala	Ala	Thr	Ala
Thr 465	Ala	Ser	Glu	His	Ser 470	Arg	Glu	Pro	Ser	Ala 475	Ala	Gly	Arg	Leu	Ser 480
Asp	Ser	Ser	Ser	Glu 485	Ala	Ser	Lys	Leu	Ser 490	Ser	Lys	Ser	Ala	Lys 495	Glu
Arg	Arg	Asn	Arg 500	Arg	Lys	Lys	Arg	Lys 505	Gln	Lys	Glu	Gln	Ser 510	Gly	Gly
Glu	Glu	Lys 51 5	Asp	Glu	Asp	Glu	Phe 520	Gln	Lys	Ser	Glu	Ser 525	Glu	Asp	Ser
Ile	Arg 530	Arg	Lys	Gly	Phe	Arg 535	Phe	Ser	Ile	Glu	Gly 540	Asn	Arg	Leu	Thr
Tyr 545	Glu	Lys	Arg	Tyr	Ser 550	Ser	Pro	His	Gln	Ser 555	Leu	Leu	Ser	Ile	Arg 560
Gly	Ser	Leu	Phe	Ser 565	Pro	Arg	Arg	Asn	Ser 570	Arg	Thr	Ser	Leu	Phe 575	Ser
Phe	Arg	Gly	Arg 580	Ala	Lys	Asp	Val	Gly 585	Ser	Glu	Asn	Asp	Phe 590	Ala	Asp

SCN1APCT1.ST25.txt

Asp Glu His Ser Thr Phe Glu Asp Asn Glu Ser Arg Asp Ser Leu
595 600 605

Phe Val Pro Arg Arg His Gly Glu Arg Arg Asn Ser Asn Leu Ser Gln 610 615 620

Thr Ser Arg Ser Ser Arg Met Leu Ala Val Phe Pro Ala Asn Gly Lys 625 630 635 640

Met His Ser Thr Val Asp Cys Asn Gly Val Val Ser Leu Val Gly Gly 645 650 655

Pro Ser Val Pro Thr Ser Pro Val Gly Gln Leu Leu Pro Glu Val Ile 660 665 670

Ile Asp Lys Pro Ala Thr Asp Asp Asn Gly Thr Thr Thr Glu Thr Glu 675 680 685

Met Arg Lys Arg Arg Ser Ser Ser Phe His Val Ser Met Asp Phe Leu 690 695 700

Glu Asp Pro Ser Gln Arg Gln Arg Ala Met Ser Ile Ala Ser Ile Leu 705 710 715 720

Thr Asn Thr Val Glu Glu Leu Glu Glu Ser Arg Gln Lys Cys Pro Pro 725 730 735

Cys Trp Tyr Lys Phe Ser Asn Ile Phe Leu Ile Trp Asp Cys Ser Pro 740 745 750

Tyr Trp Leu Lys Val Lys His Val Val Asn Leu Val Val Met Asp Pro 755 760 765

Phe Val Asp Leu Ala Ile Thr Ile Cys Ile Val Leu Asn Thr Leu Phe 770 780

Met Ala Met Glu His Tyr Pro Met Thr Asp His Phe Asn Asn Val Leu 785 790 795 800

Thr Val Gly Asn Leu Val Phe Thr Gly Ile Phe Thr Ala Glu Met Phe 805 810 815

SCN1APCT1.ST25.txt

Leu Lys Ile Ile Ala Met Asp Pro Tyr Tyr Tyr Phe Gln Glu Gly Trp 820 825 830

Asn Ile Phe Asp Gly Phe Ile Val Thr Leu Ser Leu Val Glu Leu Gly 835 840 845

Leu Ala Asn Val Glu Gly Leu Ser Val Leu Arg Ser Phe Arg Leu Leu 850 855 860

Arg Val Phe Lys Leu Ala Lys Ser Trp Pro Thr Leu Asn Met Leu Ile 865 870 875 880

Lys Ile Ile Gly Asn Ser Val Gly Ala Leu Gly Asn Leu Thr Leu Val 885 890 895

Leu Ala Ile Ile Val Phe Ile Phe Ala Val Val Gly Met Gln Leu Phe 900 905 910

Gly Lys Ser Tyr Lys Asp Cys Val Cys Lys Ile Ala Ser Asp Cys Gln 915 920 925

Leu Pro Arg Trp His Met Asn Asp Phe Phe His Ser Phe Leu Ile Val 930 935 940

Phe Arg Val Leu Cys Gly Glu Trp Ile Glu Thr Met Trp Asp Cys Met 945 950 955 960

Glu Val Ala Gly Gln Ala Met Cys Leu Thr Val Phe Met Met Val Met 965 970 975

Val Ile Gly Asn Leu Val Val Leu Asn Leu Phe Leu Ala Leu Leu Leu 980 985 990

Ser Ser Phe Ser Ala Asp Asn Leu Ala Ala Thr Asp Asp Asn Glu 995 1000 1005

Met Asn Asn Leu Gln Ile Ala Val Asp Arg Met His Lys Gly Val 1010 1015 1020

Ala Tyr Val Lys Arg Lys Ile Tyr Glu Phe Ile Gln Gln Ser Phe 1025 1030 1035

Page 15

SCN1APCT1.ST25.txt

Ile	Arg 1040	Lys	Gln	Lys	Ile	Leu 1045	Asp	Glu	Ile	Lys	Pro 1050	Leu	Asp	Asp
Leu	Asn 1055	Asn	Lys	Lys	Asp	Ser 1060	Cys	Met	Ser	Asn	ніs 1065	Thr	Thr	G1u
Ile	Gly 1070	Lys	Asp	Leu	Asp	Tyr 1075	Leu	Lys	Asp	Val	Asn 1080	Gly	Thr	Thr
Ser	Gly 1085		Gly	Thr	Gly	Ser 1090	Ser	Val	Glu	Lys	Tyr 1095	Ile	Ile	Asp
Glu	Ser 1100	Asp	Tyr	Met	Ser	Phe 1105	Ile	Asn	Asn	Pro	Ser 1110	Leu	Thr	Val
Thr	Val 1115	Pro	Ile	Ala	Val	Gly 1120	Glu	Ser	Asp	Phe	Glu 1125	Asn	Leu	Asn
Thr	Glu 1130	Asp	Phe	Ser	Ser	Glu 1135	Ser	Asp	Leu	Glu	Glu 1140	Ser	Lys	Glu
Lys	Leu 1145	Asn	Glu	Ser	Ser	Ser 1150	Ser	Ser	Glu	Gly	Ser 1155	Thr	Val	Asp
Ile	Gly 1160	Ala	Pro	Val	Glu	Glu 1165	Gln	Pro	Val	Val	Glu 1170	Pro	Glu	Glu
Thr	Leu 1175	Glu	Pro	Glu	Ala	Cys 1180	Phe	Thr	Glu	Gly	Cys 1185	Val	Gln	Arg
Phe	Lys 1190	Cys	Cys	Gln	Ile	Asn 1 1 95	Val	Glu	Glu	Gly	Arg 1200	Gly	Lys	Gln
Trp	Trp 1205	Asn	Leu	Arg	Arg	Thr 1210	Cys	Phe	Arg	Ile	Val 1215	Glu	His	Asn
Trp	Phe 1220	Glu	Thr	Phe	Ile	Val 1225	Phe	Met	Ile	Leu	Leu 1230	Ser	Ser	Gly
Ala	Leu	Ala	Phe	Glu	Asp	Ile		Ile e 16	Asp	G1n	Arg	Lys	Thr	Ile

1245

SCN1APCT1.ST25.txt 1235 1240

	123 3										1213			
Lys	Thr 1250		Leu	Glu	Tyr	Ala 1255		Lys	Val	Phe	Thr 1260		Ile	Phe
Ile	Leu 1265	Glu	Met	Leu	Leu	Lys 1270	Trp	Val	Ala	Tyr	Gly 1275	Tyr	Gln	Thr
Tyr	Phe 1280	Thr	Asn	Ala	Trp	Cys 1285		Leu	Asp	Phe	Leu 1290	Ile	Val	Asp
Val	Ser 1295	Leu	Val	Ser	Leu	Thr 1300	Ala	Asn	Ala	Leu	Gly 1305	Tyr	Ser	Glu
Leu	Gly 1310	Ala	Ile	Lys	Ser	Leu 1315		Thr	Leu	Arg	Ala 1320	Leu	Arg	Pro
Leu	Arg 1325		Leu	Ser		Phe 1330		Gly	Met	Arg	Val 1335		Val	Asn
Ala	Leu 1340	Leu	Gly	Ala	Ile	Pro 1345	Ser	Ile	Met	Asn	Val 1350	Leu	Leu	Val
Cys	Leu 1355	Ile	Phe	Trp	Leu	I1e 1360	Phe	Ser	Ile	Met	Gly 1365	Val	Asn	Leu
Phe	Ala 1370	Gly	Lys	Phe		His 1375		Ile	Asn	Thr	Thr 1380	Thr	Gly	Asp
Arg	Phe 1385	Asp	Ile	Glu	Asp	Val 1390	Asn	Asn	His	Thr	Asp 1395	Cys	Leu	Lys
Leu	Ile 1400	Glu	Arg	Asn	Glu	Thr 1405	Ala	Arg	Trp	Lys	Asn 1410	Val	Lys	Val
Asn	Phe 1415	Asp	Asn	Val	Gly	Phe 1420	Gly	Tyr	Leu	Ser	Leu 1425	Leu	Gln	Val
Ala	Thr 1430	Phe	Lys	Gly	Trp	Met 1435	Asp	Ile	Met	Tyr	Ala 1440	Ala	Val	Asp

						SCN1	APCT1	.ST2	5.tx	ct				
Ser	Arg 1445	Asn	Val	Glu	Leu						Lys 1455	Ser	Leu	Tyr
Met	Tyr 1460	Leu	Tyr	Phe	Val	Ile 1465	Phe	Ile	Ile	Phe	Gly 1470	Ser	Phe	Phe
Thr	Leu 1475	Asn	Leu	Phe	Ile	Gly 1480	Val	Ile	I1e	Asp	Asn 1485	Phe	Asn	Gln
G1n	Lys 1490	Lys	Lys	Phe	Gly	Gly 1495	Gln	Asp	Ile	Phe	Met 1500	Thr	Glu	Glu
Gln	Lys 1505		Tyr	Tyr	Asn	A1a 1510		Lys	Lys	Leu	Gly 1515		Lys	Lys
Pro	Gln 1520	Lys	Pro	Ile	Pro	Arg 1525	Pro	Gly	Asn	Lys	Phe 1530	Gln	Gly	Met
Val	Phe 1535	Asp	Phe	Val	Thr	Arg 1540	Gln	Val	Phe	Asp	Ile 1545	Ser	Ile	Met
Ile	Leu 1550	Ile	Cys	Leu	Asn	Met 1555	Val	Thr	Met	Met	Val 1560	Glu	Thr	Asp
Asp	Gln 1565	Ser	Glu	Tyr	Val	Thr 1570	Thr	Ile	Leu	Ser	Arg 1575	Ile	Asn	Leu
Val	Phe 1580	Ile	Val	Leu	Phe	T hr 1585	Gly	Glu	Cys	Val	Leu 1590	Lys	Leu	Ile
Ser	Leu 1595	Arg	His	Tyr	Tyr	Phe 1600	Thr	Ile	Gly	Trp	Asn 1605	Ile	Phe	Asp
Phe	Val 1610	Val	Val	Ile	Leu	Ser 1615	Ile	Val	Gly	Met	Phe 1620	Leu	Ala	Glu
Leu	Ile 1625	Glu	Lys	Tyr	Phe	Val 1630	Ser	Pro	Thr	Leu	Phe 1635	Arg	Val	Ile
Arg	Leu 1640	Ala	Arg	Ile	Gly	Arg 1645	Ile	Leu	Arg	Leu	Ile 1650	Lys	Gly	Ala

SCN1APCT1.ST25.txt

Lys	Gly 1655	Ile	Arg	Thr	Leu	Leu 1660	Phe	Ala	Leu	Met	Met 1665	Ser	Leu	Pro
Ala	Leu 1670	Phe	Asn	Ile	Gly	Leu 1675	Leu	Leu	Phe	Leu	Val 1680	Met	Phe	Ile
Tyr	A1a 1685	I1e	Phe	Gly	Met	Ser 1690	Asn	Phe	Ala	Tyr	Val 1695	Lys	Arg	Glu
Val	G1y 1700	I1e	Asp	Asp	Met	Phe 1705	Asn	Phe	Glu	Thr	Phe 1710	G1y	Asn	Ser
Met	Ile 1715	Cys	Leu	Phe	Gln	Ile 1720	Thr	Thr	Ser	Ala	G1y 1725	Trp	Asp	G1y
Leu	Leu 1730	Ala	Pro	Ile	Leu	Asn 1735	Ser	Lys	Pro	Pro	Asp 1740	Cys	Asp	Pro
Asn	Lys 1745		Asn	Pro	Gly	Ser 1750	Ser	Val	Lys	Gly	Asp 1755		Gly	Asn
Pro	Ser 1760	Va1	Gly	Ile	Phe	Phe 1765	Phe	Val	Ser	Tyr	Ile 1770	Ile	Ile	Ser
Phe	Leu 1775	Va1	Val	Val	Asn	Met 1780	Tyr	Ile	Ala	Val	Ile 1785	Leu	Glu	Asn
Phe	Ser 1790	Va1	Ala	Thr	Glu	Glu 1795	Ser	Ala	Glu	Pro	Leu 1800	Ser	G1u	Asp
Asp	Phe 1805	G1u	Met	Phe	Tyr	Glu 1810	Val	Trp	Glu	Lys	Phe 1815	Asp	Pro	Asp
A1a	Thr 1820	G1n	Phe	Met	Glu	Phe 1825	Glu	Lys	Leu	Ser	G1n 1830	Phe	Ala	Ala
Ala	Leu 1835	Glu	Pro	Pro	Leu	Asn 1840	Leu	Pro	Gln	Pro	Asn 1845	Lys	Leu	G1n
Leu	Ile 1850	Ala	Met	Asp	Leu	Pro 1855		Val e 19	Ser	Gly	Asp 1860	Arg	I1e	His

SCN1APCT1.ST25.txt

Cys	Leu 1865	Asp	Ile	Leu	Phe	Ala 1870	Phe	Thr	Lys	Arg	Val 1875	Leu	Gly	Glu
Ser	Gly 1880	Glu	Met	Asp	Ala	Leu 1885	Arg	Ile	Gln	Met	Glu 1890	Glu	Arg	Phe

- Met Ala Ser Asn Pro Ser Lys Val Ser Tyr Gln Pro Ile Thr Thr 1895 1900 1905
- Thr Leu Lys Arg Lys Gln Glu Glu Val Ser Ala Val Ile Ile Gln 1910 1915 1920
- Arg Ala Tyr Arg Arg His Leu Leu Lys Arg Thr Val Lys Gln Ala 1925 1930 1935
- Ser Phe Thr Tyr Asn Lys Asn Lys Ile Lys Gly Gly Ala Asn Leu 1940 1945 1950
- Leu Ile Lys Glu Asp Met Ile Ile Asp Arg Ile Asn Glu Asn Ser 1955 1960 1965
- Ile Thr Glu Lys Thr Asp Leu Thr Met Ser Thr Ala Ala Cys Pro 1970 1975 1980
- Pro Ser Tyr Asp Arg Val Thr Lys Pro Ile Val Glu Lys His Glu 1985 1990 1995
- Gln Glu Gly Lys Asp Glu Lys Ala Lys Gly Lys 2000 2005

<210> 3<211> 8381<212> DNA<213> Homo sapiens<400> 3
atactgcaga ggtctctggt gcatgtgtgt atgtgtgcgt ttgtgtgtgt ttgtgtgtct
60

gtgtgttctg ccccagtgag actgcagccc ttgtaaatac tttgacacct tttgcaagaa 120

ggaatctgaa caattgcaac tgaaggcaca ttgttatcat ctcgtctttg ggtgatgctg
180

SCN1APCT1.ST25.txt

ttcctcactg cagatggata attttccttt taatcaggaa tttcatatgc agaataaatg 240

gtaattaaaa tgtgcaggat gacaagatgg agcaaacagt gcttgtacca ccaggacctg

acagetteaa ettetteace agagaatete ttgeggetat tgaaagaege attgeagaag

360

aaaaggcaaa gaatcccaaa ccagacaaaa aagatgacga cgaaaatggc ccaaagccaa 420

atagtgactt ggaagctgga aagaaccttc catttattta tggagacatt cctccagaga 480

tggtgtcaga gcccctggag gacctggacc cctactatat caataagaaa acttttatag
540

tattgaataa attgaaggcc atcttccggt tcagtgccac ctctgccctg tacattttaa 600

ctcccttcaa tcctcttagg aaaatagcta ttaagatttt ggtacattca ttattcagca 660

tgctaattat gtgcactatt ttgacaaact gtgtgtttat gacaatgagt aaccetectg
720

attggacaaa gaatgtagaa tacaccttca caggaatata tacttttgaa tcacttataa 780

aaattattgc aaggggattc tgtttagaag attttacttt ccttcgggat ccatggaact 840

ggctcgattt cactgtcatt acatttgcgt acgtcacaga gtttgtggac ctgggcaatg
900

tctcggcatt gagaacattc agagttctcc gagcattgaa gacgatttca gtcattccag
960

gcctgaaaac cattgtggga gccctgatcc agtctgtgaa gaagctctca gatgtaatga 1020

SCN1APCT1.ST25.txt

tcctgactgt gttctgtctg agegtatttg ctctaattgg gctgcagctg ttcatgggca
1080

acctgaggaa taaatgtata caatggcctc ccaccaatgc ttccttggag gaacatagta 1140

tagaaaagaa tataactgtg aattataatg gtacacttat aaatgaaact gtctttgagt 1200

ttgactggaa gtcatatatt caagattcaa gatatcatta tttcctggag ggttttttag
1260

atgcactact atgtggaaat agctctgatg caggccaatg tccagaggga tatatgtgtg
1320

tgaaagctgg tagaaatccc aattatggct acacaagctt tgataccttc agttgggctt 1380

ttttgtcctt gtttcgacta atgactcagg acttctggga aaatctttat caactgacat 1440

tacgtgctgc tgggaaaacg tacatgatat tttttgtatt ggtcattttc ttgggctcat
1500

tctacctaat aaatttgatc ctggctgtgg tggccatggc ctacgaggaa cagaatcagg 1560

ccaccttgga agaagcagaa cagaaagagg ccgaatttca gcagatgatt gaacagctta
1620

aaaagcaaca ggaggcagct cagcaggcag caacggcaac tgcctcagaa cattccagag

agcccagtgc agcaggcagg ctctcagaca gctcatctga agcctctaag ttgagttcca

gggaagagaa agatgaggat gaattccaaa aatctgaatc tgaggacagc atcaggagga

SCN1APCT1.ST25.txt

1860

aaggttttcg cttctccatt gaagggaacc gattgacata tgaaaagagg tactcctccc

1920

cacaccagtc tttgttgagc atccgtggct ccctattttc accaaggcga aatagcagaa

1980

caageetttt cagetttaga gggegageaa aggatgtggg atetgagaae gaettegeag

2040

atgatgagca cagcaccttt gaggataacg agagccgtag agattccttg tttgtgcccc

2100

gacgacacgg agagagacgc aacagcaacc tgagtcagac cagtaggtca tcccggatgc

2160

tggcagtgtt tccagcgaat gggaagatgc acagcactgt ggattgcaat ggtgtggttt

2220

ccttqqttqq tqqaccttca qttcctacat cgcctqttqq acagcttctq ccaqaqqtqa

2280

taatagataa gccagctact gatgacaatg gaacaaccac tgaaactgaa atgagaaaga

2340

gaaggtcaag ttctttccac gtttccatgg actttctaga agatccttcc caaaggcaac

2400

gaqcaatqag tatagccagc attctaacaa atacagtaga agaacttgaa gaatccaggc

2460

agaaatgccc accctgttgg tataaatttt ccaacatatt cttaatctgg gactgttctc

2520

catattggtt aaaagtgaaa catgttgtca acctggttgt gatggaccca tttgttgacc

2580

tggccatcac catctgtatt gtcttaaata ctcttttcat ggccatggag cactatccaa

2640

tgacggacca tttcaataat gtgcttacag taggaaactt ggttttcact gggatcttta

Page 23

SCN1APCT1.ST25.txt

2700

cagcagaaat gtttctgaaa attattgcca tggatcctta ctattatttc caagaaggct 2760

ggaatatett tgaeggtttt attgtgaege ttageetggt agaaettgga etegeeaatg 2820

tggaaggatt atctgttctc cgttcatttc gattgctgcg agttttcaag ttggcaaaat 2880

cttggccaac gttaaatatg ctaataaaga tcatcggcaa ttccgtgggg gctctgggaa 2940

atttaaccct cgtcttggcc atcatcgtct tcatttttgc cgtggtcggc atgcagctct 3000

ttggtaaaag ctacaaagat tgtgtctgca agatcgccag tgattgtcaa ctcccacgct 3060

ggcacatgaa tgacttcttc cactccttcc tgattgtgtt ccgcgtgctg tgtggggagt 3120

ggatagagac catgtgggac tgtatggagg ttgctggtca agccatgtgc cttactgtct 3180

tcatgatggt catggtgatt ggaaacctag tggtcctgaa tctctttctg gccttgcttc 3240

tgagctcatt tagtgcagac aaccttgcag ccactgatga tgataatgaa atgaataatc 3300

tccaaattgc tgtggatagg atgcacaaag gagtagctta tgtgaaaaga aaaatatatg 3360

aatttattca acagtccttc attaggaaac aaaagatttt agatgaaatt aaaccacttg 3420

atgatctaaa caacaagaaa gacagttgta tgtccaatca tacaacagaa attgggaaag 3480

SCN1APCT1_ST25.txt

atcttgacta tcttaaagat gtaaatggaa ctacaagtgg tataggaact ggcagcagtg 3540

ttgaaaaata cattattgat gaaagtgatt acatgtcatt cataaacaac cccagtctta 3600

ctgtgactgt accaattgct gtaggagaat ctgactttga aaatttaaac acggaagact 3660

ttagtagtga atcggatctg gaagaaagca aagagaaact gaatgaaagc agtagctcat 3720

cagaaggtag cactgtggac atcggcgcac ctgtagaaga acagcccgta gtggaacctg 3780

aagaaactct tgaaccagaa gcttgtttca ctgaaggctg tgtacaaaga ttcaagtgtt 3840

gtcaaatcaa tgtggaagaa ggcagaggaa aacaatggtg gaacctgaga aggacgtgtt 3900

tccgaatagt tgaacataac tggtttgaga ccttcattgt tttcatgatt ctccttagta 3960

gtggtgctct ggcatttgaa gatatatata ttgatcagcg aaagacgatt aagacgatgt 4020

tggaatatgc tgacaaggtt ttcacttaca ttttcattct ggaaatgctt ctaaaatggg 4080

tggcatatgg ctatcaaaca tatttcacca atgcctggtg ttggctggac ttcttaattg 4140

ttgatgtttc attggtcagt ttaacagcaa atgccttggg ttactcagaa cttggagcca 4200

tcaaatctct caggacacta agagctctga gacctctaag agccttatct cgatttgaag 4260

ggatgagggt ggttgtgaat gcccttttag gagcaattcc atccatcatg aatgtgcttc 4320

SCN1APCT1.ST25.txt

tgctttgtct tatattctgg ctaattttca gcatcatggg cgtaaatttg tttgctggca 4380

aattctacca ctgtattaac accacaactg gtgacaggtt tgacatcgaa gacgtgaata 4440

atcatactga ttgcctaaaa ctaatagaaa gaaatgagac tgctcgatgg aaaaatgtga 4500

aagtaaactt tgataatgta ggatttgggt atctctcttt gcttcaagtt gccacattca 4560

aaggatggat ggatataatg tatgcagcag ttgattccag aaatgtggaa ctccagccta 4620

agtatgaaaa aagtctgtac atgtatcttt actttgttat tttcatcatc tttgggtcct 4680

tcttcacctt gaacctgttt attggtgtca tcatagataa tttcaaccag cagaaaaaga 4740

agtttggagg tcaagacatc tttatgacag aagaacagaa gaaatactat aatgcaatga 4800

aaaaattagg atcgaaaaaa ccgcaaaagc ctatacctcg accaggaaac aaatttcaag 4860

gaatggtctt tgacttcgta accagacaag tttttgacat aagcatcatg attctcatct 4920

gtcttaacat ggtcacaatg atggtggaaa cagatgacca gagtgaatat gtgactacca 4980

ttttgtcacg catcaatctg gtgttcattg tgctatttac tggagagtgt gtactgaaac 5040

tcatctctct acgccattat tattttacca ttggatggaa tatttttgat tttgtggttg 5100

tcattctctc cattgtaggt atgtttcttg ccgagctgat agaaaagtat ttcgtgtccc

SCN1APCT1.ST25.txt

5160

ctaccetgtt eegagtgate egtettgeta ggattggeeg aateetaegt etgateaaag 5220

gagcaaaggg gatccgcacg ctgctctttg ctttgatgat gtcccttcct gcgttgttta 5280

acateggeet ectaetette etagteatgt teatetaege eatetttggg atgteeaact 5340

ttgcctatgt taagagggaa gttgggatcg atgacatgtt caactttgag acctttggca 5400

acagcatgat ctgcctattc caaattacaa cctctgctgg ctgggatgga ttgctagcac 5460

ccattctcaa cagtaagcca cccgactgtg accctaataa agttaaccct ggaagctcag 5520

ttaagggaga ctgtgggaac ccatctgttg gaattttctt ttttgtcagt tacatcatca
5580

tatccttcct ggttgtggtg aacatgtaca tcgcggtcat cctggagaac ttcagtgttg 5640

ctactgaaga aagtgcagag cctctgagtg aggatgactt tgagatgttc tatgaggttt 5700

gggagaagtt tgatcccgat gcaactcagt tcatggaatt tgaaaaatta tctcagtttg 5760

cagctgcgct tgaaccgcct ctcaatctgc cacaaccaaa caaactccag ctcattgcca 5820

tggatttgcc catggtgagt ggtgaccgga tccactgtct tgatatctta tttgctttta
5880

caaagcgggt tctaggagag agtggagaga tggatgctct acgaatacag atggaagagc 5940

gattcatggc ttccaatcct tccaaggtct cctatcagcc aatcactact actttaaaac Page 27

SCN1APCT1.ST25.txt

6000

gaaaacaaga ggaagtatct gctgtcatta ttcagcgtgc ttacagacgc caccttttaa 6060

agcgaactgt aaaacaagct tcctttacgt acaataaaaa caaaatcaaa ggtggggcta 6120

atcttcttat aaaagaagac atgataattg acagaataaa tgaaaactct attacagaaa 6180

aaactgatct gaccatgtcc actgcagctt gtccaccttc ctatgaccgg gtgacaaagc 6240

caattgtgga aaacatgag caagaaggca aagatgaaaa agccaaaggg aaataaatga 6300

aaataaataa aaataattgg gtgacaaatt gtttacagcc tgtgaaggtg atgtattttt 6360

atcaacagga ctcctttagg aggtcaatgc caaactgact gtttttacac aaatctcctt 6420

aaggtcagtg cctacaataa gacagtgacc ccttgtcagc aaactgtgac tctgtgtaaa 6480

ggggagatga cettgacagg aggttactgt tetcactace agetgacaet getgaagata 6540

agatgcacaa tggctagtca gactgtaggg accagtttca aggggtgcaa acctgtgatt 6600

ttggggttgt ttaacatgaa acactttagt gtagtaattg tatccactgt ttgcatttca 6660

actgccacat ttgtcacatt tttatggaat ctgttagtgg attcatcttt ttgttaatcc 6720

atgtgtttat tatatgtgac tatttttgta aacgaagttt ctgttgagaa ataggctaag 6780

SCN1APCT1.ST25.txt

gacetetata acaggtatge cacetggggg gtatggcaac cacatggeee teccagetae 6840

acaaagtcgt ggtttgcatg agggcatgct gcacttagag atcatgcatg agaaaaagtc 6900

acaagaaaaa caaattotta aatttoacca tatttotggg aggggtaatt gggtgataag 6960

tggaggtgct ttgttgatct tgttttgcga aatccagccc ctagaccaag tagattattt 7020

gtgggtaggc cagtaaatct tagcaggtgc aaacttcatt caaatgtttg gagtcataaa 7080

tgttatgttt ctttttgttg tattaaaaaa aaaacctgaa tagtgaatat tgcccctcac 7140

cctccaccgc cagaagactg aattgaccaa aattactctt tataaatttc tgctttttcc
7200

tgcactttgt ttagccatct ttgggctctc agcaaggttg acactgtata tgttaatgaa 7260

atgctattta ttatgtaaat agtcatttta ccctgtggtg cacgtttgag caaacaaata 7320

atgacctaag cacagtattt attgcatcaa atatgtacca caagaaatgt agagtgcaag 7380

ctttacacag gtaataaaat gtattctgta ccatttatag atagtttgga tgctatcaat 7440

gcatgtttat attaccatgc tgctgtatct ggtttctctc actgctcaga atctcattta 7500

tgagaaacca tatgtcagtg gtaaagtcaa ggaaattgtt caacagatct catttattta
7560

agtcattaag caatagtttg cagcacttta acagcttttt ggttatttt acattttaag

SCN1APCT1.ST25.txt

tggataacat atggtatata gccagactgt acagacatgt ttaaaaaaac acactgctta 7680

acctattaaa tatgtgttta gaattttata agcaaatata aatactgtaa aaagtcactt 7740

tattttattt ttcagcatta tgtacataaa tatgaagagg aaattatctt caggttgata 7800

tcacaatcac ttttcttact ttctgtccat agtactttt catgaaagaa atttgctaaa 7860

taagacatga aaacaagact gggtagttgt agatttctgc tttttaaatt acatttgcta 7920

attttagatt atttcacaat tttaaggagc aaaataggtt cacgattcat atccaaatta 7980

tgctttgcaa ttggaaaagg gtttaaaatt ttatttatat ttctggtagt acctgtacta 8040

actgaattga aggtagtgct tatgttattt ttgttctttt tttctgactt cggtttatgt 8100

tttcatttct ttggagtaat gctgctctag attgttctaa atagaatgtg ggcttcataa 8160

ttttttttc cacaaaaaca gagtagtcaa cttatatagt caattacatc aggacatttt 8220

gtgtttctta cagaagcaaa ccataggctc ctcttttcct taaaactact tagataaact

gtattcgtga actgcatgct ggaaaatgct actattatgc taaataatgc taaccaacat 8340

ttaaaatgtg caaaactaat aaagattaca tttttattt t 8381

SCN1APCT1.ST25.txt

Met Glu Gln Thr Val Leu Val Pro Pro Gly Pro Asp Ser Phe Asn Phe 1 5 10 15

Phe Thr Arg Glu Ser Leu Ala Ala Ile Glu Arg Arg Ile Ala Glu Glu 20 25 30

Lys Ala Lys Asn Pro Lys Pro Asp Lys Lys Asp Asp Asp Glu Asn Gly 35 40 45

Pro Lys Pro Asn Ser Asp Leu Glu Ala Gly Lys Asn Leu Pro Phe Ile 50 55 60

Tyr Gly Asp Ile Pro Pro Glu Met Val Ser Glu Pro Leu Glu Asp Leu 65 70 75 80

Asp Pro Tyr Tyr Ile Asn Lys Lys Thr Phe Ile Val Leu Asn Lys Leu 85 90 95

Lys Ala Ile Phe Arg Phe Ser Ala Thr Ser Ala Leu Tyr Ile Leu Thr 100 105 110

Pro Phe Asn Pro Leu Arg Lys Ile Ala Ile Lys Ile Leu Val His Ser 115 120 125

Leu Phe Ser Met Leu Ile Met Cys Thr Ile Leu Thr Asn Cys Val Phe 130 135 140

Met Thr Met Ser Asn Pro Pro Asp Trp Thr Lys Asn Val Glu Tyr Thr 145 150 155 160

Phe Thr Gly Ile Tyr Thr Phe Glu Ser Leu Ile Lys Ile Ile Ala Arg 165 170 175

Gly Phe Cys Leu Glu Asp Phe Thr Phe Leu Arg Asp Pro Trp Asn Trp 180 185 190

Leu Asp Phe Thr Val Ile Thr Phe Ala Tyr Val Thr Glu Phe Val Asp 195 200 205

Leu Gly Asn Val Ser Ala Leu Arg Thr Phe Arg Val Leu Arg Ala Leu 210 215 220

Page 31

SCN1APCT1.ST25.txt

Lys 225	Thr	Ile	Ser	Val	Ile 230	Pro	Gly	Leu	Lys	Thr 235	Ile	Val	Gly	Ala	Leu 240
Ile	Gln	Ser	Val	Lys 245	Lys	Leu	Ser	Asp	Val 250	Met	Ile	Leu	Thr	Val 255	Phe
Cys	Leu	Ser	Val 260	Phe	Ala	Leu	Ile	Gly 265	Leu	Gln	Leu	Phe	Met 270	Gly	Asn
Leu	Arg	Asn 275	Lys	Cys	Ile	Gln	Trp 280	Pro	Pro	Thr	Asn	Ala 285	Ser	Leu	Glu
Glu	His 290	Ser	Ile	Glu	Lys	Asn 295	Ile	Thr	Val	Asn	Tyr 300	Asn	Gly	Thr	Leu
Ile 305	Asn	Glu	Thr	Val	Phe 310	Glu	Phe	Asp	Trp	Lys 315	Ser	Tyr	Ile	Gln	Asp 320
Ser	Arg	Tyr	His	Tyr 325	Phe	Leu	Glu	Gly	Phe 330	Leu	Asp	Ala	Leu	Leu 335	Cys
Gly	Asn	Ser	Ser 340	Asp	Ala	Gly	Gln	Cys 345	Pro	Glu	Gly	Tyr	Met 350	Cys	Val
Lys	Ala	Gly 355	Arg	Asn	Pro	Asn	Туг 360	Gly	Tyr	Thr	Ser	Phe 365	Asp	Thr	Phe
Ser	Trp 370	Ala	Phe	Leu	Ser	Leu 375	Phe	Arg	Leu	Met	Thr 380	Gln	Asp	Phe	Trp
Glu 385	Asn	Leu	Tyr	Gln	Leu 390	Thr	Leu	Arg	Ala	Ala 395	Gly	Lys	Thr	Tyr	Met 400
Ile	Phe	Phe	Val	Leu 405	Val	Ile	Phe	Leu	Gly 410	Ser	Phe	Tyr	Leu	Ile 415	Asn
Leu	Ile	Leu	Ala 420	Val	Val	Ala	Met	Ala 425	Tyr	Glu	Glu	Gln	Asn 430	Gln	Ala
Thr	Leu	Glu	Glu	Ala	Glu	Gln	_	Glu age		Glu	Phe	Gln	Gln	Met	Ile

WO 02/50096	PCT/AU01/01648

445

SCN1APCT1.ST25.txt 435 440

Glu	Gln 450	Leu	Lys	Lys	Gln	Gln 455	Glu	Ala	Ala	Gln	Gln 460	Ala	Ala	Thr	Ala
Thr 465	Ala	Ser	Glu	His	Ser 470	Arg	Glu	Pro	Ser	Ala 475	Ala	Gly	Arg	Leu	Ser 480
Asp	Ser	Ser	Ser	Glu 485	Ala	Ser	Lys	Leu	Ser 490	Ser	Lys	Ser	Ala	Lys 495	Glu
Arg	Arg	Asn	Arg 500	Arg	Lys	Lys	Arg	Lys 505	Gln	Lys	Glu	Gln	Ser 510	Gly	Gly
Glu	Glu	Lys 515	Asp	Glu	Asp	Glu	Phe 520	Gln	Lys	Ser	Glu	Ser 525	Glu	Asp	Ser
Ile	Arg 530	Arg	Lys	Gly	Phe	Arg 535	Phe	Ser	Ile	Glu	Gly 540	Asn	Arg	Leu	Thr
Tyr 545	Glu	Lys	Arg	Tyr	Ser 550	Ser	Pro	His	Gln	Ser 555	Leu	Leu	Ser	Ile	Arg 560
Gly	Ser	Leu	Phe	Ser 565	Pro	Arg	Arg	Asn	Ser 570	Arg	Thr	Ser	Leu	Phe 575	Ser
Phe	Arg	Gly	Arg 580	Ala	Lys	Asp	Val	Gly 585	Ser	Glu	Asn	Asp	Phe 590	Ala	Asp
Asp	Glu	His 595	Ser	Thr	Phe	Glu	Asp 600	Asn	Glu	Ser	Arg	Arg 605	Asp	Ser	Leu
Phe	Val 610	Pro	Arg	Arg	His	Gly 615	Glu	Arg	Arg	Asn	Ser 620	Asn	Leu	Ser	Gln
Thr 625	Ser	Arg	Ser	Ser	Arg 630	Met	Leu	Ala	Val	Phe 635	Pro	Ala	Asn	Gly	Lys 640
Met	His	Ser	Thr	Val 645	Asp	Cys	Asn	Gly	Val 650	Val	Ser	Leu	Val	Gly 655	Gly

SCN1APCT1.ST25.txt

Pro Ser Val Pro Thr Ser Pro Val Gly Gln Leu Leu Pro Glu Val Ile 660 665 670

Ile Asp Lys Pro Ala Thr Asp Asp Asn Gly Thr Thr Thr Glu Thr Glu 675 680 685

Met Arg Lys Arg Arg Ser Ser Ser Phe His Val Ser Met Asp Phe Leu 690 695 700

Glu Asp Pro Ser Gln Arg Gln Arg Ala Met Ser Ile Ala Ser Ile Leu 705 710 715 720

Thr Asn Thr Val Glu Glu Leu Glu Glu Ser Arg Gln Lys Cys Pro Pro 725 730 735

Cys Trp Tyr Lys Phe Ser Asn Ile Phe Leu Ile Trp Asp Cys Ser Pro 740 745 750

Tyr Trp Leu Lys Val Lys His Val Val Asn Leu Val Val Met Asp Pro 755 760 765

Phe Val Asp Leu Ala Ile Thr Ile Cys Ile Val Leu Asn Thr Leu Phe 770 775 780

Met Ala Met Glu His Tyr Pro Met Thr Asp His Phe Asn Asn Val Leu 785 790 795 800

Thr Val Gly Asn Leu Val Phe Thr Gly Ile Phe Thr Ala Glu Met Phe 805 810 815

Leu Lys Ile Ile Ala Met Asp Pro Tyr Tyr Tyr Phe Gln Glu Gly Trp 820 825 830

Asn Ile Phe Asp Gly Phe Ile Val Thr Leu Ser Leu Val Glu Leu Gly 835 840 845

Leu Ala Asn Val Glu Gly Leu Ser Val Leu Arg Ser Phe Arg Leu Leu 850 855 860

Arg Val Phe Lys Leu Ala Lys Ser Trp Pro Thr Leu Asn Met Leu Ile 865 870 875 880

SCN1APCT1.ST25.txt

Lys Ile Ile Gly Asn Ser Val Gly Ala Leu Gly Asn Leu Thr Leu Val 885 890 895

Leu Ala Ile Ile Val Phe Ile Phe Ala Val Val Gly Met Gln Leu Phe 900 905 910

Gly Lys Ser Tyr Lys Asp Cys Val Cys Lys Ile Ala Ser Asp Cys Gln 915 920 925

Leu Pro Arg Trp His Met Asn Asp Phe Phe His Ser Phe Leu Ile Val 930 935 940

Phe Arg Val Leu Cys Gly Glu Trp Ile Glu Thr Met Trp Asp Cys Met 945 950 955 960

Glu Val Ala Gly Gln Ala Met Cys Leu Thr Val Phe Met Met Val Met 965 970 975

Val Ile Gly Asn Leu Val Val Leu Asn Leu Phe Leu Ala Leu Leu 980 985 990

Ser Ser Phe Ser Ala Asp Asn Leu Ala Ala Thr Asp Asp Asp Asn Glu 995 1000 1005

Met Asn Asn Leu Gln Ile Ala Val Asp Arg Met His Lys Gly Val 1010 1015 1020

Ala Tyr Val Lys Arg Lys Ile Tyr Glu Phe Ile Gln Gln Ser Phe 1025 1030 1035

Ile Arg Lys Gln Lys Ile Leu Asp Glu Ile Lys Pro Leu Asp Asp 1040 1045 1050

Leu Asn Asn Lys Lys Asp Ser Cys Met Ser Asn His Thr Thr Glu 1055 1060 1065

Ile Gly Lys Asp Leu Asp Tyr Leu Lys Asp Val Asn Gly Thr Thr 1070 1080

Ser Gly Ile Gly Thr Gly Ser Ser Val Glu Lys Tyr Ile Ile Asp 1085 1090 1095

Page 35

SCN1APCT1.ST25.txt

G1u	Ser 1100		Tyr	Met	Ser	Phe 1105	Ile	Asn	Asn	Pro	Ser 1110	Leu	Thr	Val
Thr	Val 1115	Pro	I1e	A1a	Val	G1y 1120	G1u	Ser	Asp	Phe	G1u 1125	Asn	Leu	Asn
Thr	G1u 1130		Phe	Ser	Ser	G1u 1135		Asp	Leu	G1u	G1u 1140		Lys	Glu
Lys	Leu 1145	Asn	G1u	Ser	Ser	Ser 1150	Ser	Ser	Glu	Gly	Ser 1155	Thr	Va1	Asp
Ile	G1y 1160	Ala	Pro	Va1	G1u	Glu 1165	G1n	Pro	Va1	Val	Glu 1170	Pro	Glu	Glu
Thr	Leu 1175		Pro	Glu	Ala	Cys 1180		Thr	Glu	Gly	Cys 1185	Val	Gln	Arg
Phe	Lys 1190		Cys	Gln	Ile	Asn 1195	Va1	Glu	G1u	G1y	Arg 1200	Ġly	Lys	Gln
Trp	Trp 1205		Leu	Arg	Arg	Thr 1210		Phe	Arg	Ile	Va1 1215	Glu	His	Asn
Trp	Phe 1220	Glu	Thr	Phe	Ile	Val 1225	Phe	Met	I1e	Leu	Leu 1230	Ser	Ser	Gly
Ala	Leu 1235	Ala	Phe	Glu	Asp	I1e 1240	Tyr	Ile	Asp	Gln	Arg 1245	Lys	Thr	Ile
Lys	Thr 1250	Met	Leu	Glu	Tyr	Ala 1255	Asp	Lys	Val	Phe	Thr 1260	Tyr	Ile	Phe
Ile	Leu 1265	G1u	Met	Leu	Leu	Lys 1270	Trp	Val	Ala	Tyr	G1y 1275	Tyr	G1n	Thr
Tyr	Phe 1280	Thr	Asn	Ala	Trp	Cys 1285	Trp	Leu	Asp	Phe	Leu 1290	I1e	Val	Asp
Va1	Ser	Leu	Va1	Ser	Leu	Thr		Asn e 36	Ala	Leu	Gly	Tyr	Ser	Glu

1305

SCN1APCT1.ST25.txt 1295 1300

	1275					1300					1303			
Leu	Gly 1310		Ile	Lys	Ser	Leu 1315		Thr	Leu	Arg	Ala 1320		Arg	Pro
Leu	Arg 1325	Ala	Leu	Ser	Arg	Phe 1330	Glu	Gly	Met	Arg	Val 1335	Val	Val	Asn
Ala	Leu 1340	Leu	Gly	Ala	Ile	Pro 1345	Ser	Ile	Met	Asn	Val 1350	Leu	Leu	Leu
Cys	Leu 1355		Phe	Trp	Leu	Ile 1360		Ser	Ile	Met	Gly 1365	Val	Asn	Leu
Phe	Ala 1370	Gly	Lys	Phe	Tyr	His 1375		Ile	Asn	Thr	Thr 1380	Thr	Gly	Asp
Arg	Phe 1385		Ile	Glu		Val 1390		Asn	His	Thr	Asp 1395		Leu	Lys
Leu	Ile 1400	Glu	Arg	Asn	Glu	Thr 1405	Ala	Arg	Trp	Lys	Asn 1410	Val	Lys	Val
Asn	Phe 1415	Asp	Asn	Val	Gly	Phe 1420	Gly	Tyr	Leu	Ser	Leu 1425	Leu	Gln	Val
Ala	Thr 1430	Phe	Lys	Gly	Trp	Met 1435	Asp	Ile	Met	Tyr	Ala 1440	Ala	Val	Asp
Ser	Arg 1445	Asn	Val	Glu	Leu	Gln 1450	Pro	Lys	Tyr	Glu	Lys 1455	Ser	Leu	Туг
Met	Tyr 1460	Leu	Tyr	Phe	Val	Ile 1465	Phe	Ile	Ile	Phe	Gly 1470	Ser	Phe	Phe
Thr	Leu 1475	Asn	Leu	Phe	Ile	Gly 1480	Val	Ile	Ile	Asp	Asn 1485	Phe	Asn	Gln
Gln	Lys 1490	Lys	Lys	Phe	Gly	Gly 1495	Gln	Asp	Ile	Phe	Met 1500	Thr	Glu	Glu

						SCN1	APCT1	.ST2	25.tx	ct				
Gln	Lys 1505	Lys	Tyr	Tyr	Asn						Gly 1515	Ser	Lys	Lys
Pro	Gln 1520	Lys	Pro	Ile	Pro	Arg 1525	Pro	Gly	Asn	Lys	Phe 1530	Gln	Gly	Met
Val	Phe 1535	_	Phe	Val	Thr	Arg 1540		Val	Phe	Asp	Ile 1545	Ser	Ile	Met
Ile	Leu 1550	Ile	Cys	Leu	Asn	Met 1555	Val	Thr	Met	Met	Val 1560	Glu	Thr	Asp
Asp	Gln 1565	Ser	Glu	Tyr	Va1	Thr 1570		Ile	Leu	Ser	Arg 1575		Asn	Leu
Val	Phe 1580	Ile	Val	Leu	Phe	Thr 1585	Gly	Glu	Cys	Va1	Leu 1590	Lys	Leu	Ile
Ser	Leu 1595	Arg	His	Tyr	Tyr	Phe 1600	Thr	Ile	Gly	Trp	Asn 1605	Ile	Phe	Asp
Phe	Val 1610	Val	Val	Ile	Leu	Ser 1615	Ile	Val	Gly	Met	Phe 1620	Leu	Ala	Glu
Leu	Ile 1625	Glu	Lys	Tyr	Phe	Val 1630	Ser	Pro	Thr	Leu	Phe 1635	Arg	Val	Ile
Arg	Leu 1640		Arg	Ile	Gly	Arg 1645		Leu	Arg	Leu	Ile 1650	Lys	Gly	Ala
Lys	Gly 1655	Ile	Arg	Thr	Leu	Leu 1660	Phe	Ala	Leu	Met	Met 1665	Ser	Leu	Pro
Ala	Leu 1670	Phe	Asn	Ile	Gly	Leu 1675	Leu	Leu	Phe	Leu	Val 1680	Met	Phe	Ile
Tyr	Ala 1685	Ile	Phe	Gly	Met	Ser 1690	Asn	Phe	Ala	Tyr	Val 1695	Lys	Arg	Glu
Val	Gly 1700	Ile	Asp	Asp	Met	Phe 1705	Asn	Phe	Glu	Thr	Phe 1710	Gly	Asn	Ser

SCN1APCT1.ST25.txt

Met Ile Cys Leu Phe Gln Ile Thr Thr Ser Ala Gly Trp Asp Gly Leu Leu Ala Pro Ile Leu Asn Ser Lys Pro Pro Asp Cys Asp Pro Asn Lys Val Asn Pro Gly Ser Ser Val Lys Gly Asp Cys Gly Asn 1745 1750 1755 Pro Ser Val Gly Ile Phe Phe Phe Val Ser Tyr Ile Ile Ile Ser Phe Leu Val Val Val Asn Met Tyr Ile Ala Val Ile Leu Glu Asn Phe Ser Val Ala Thr Glu Glu Ser Ala Glu Pro Leu Ser Glu Asp Asp Phe Glu Met Phe Tyr Glu Val Trp Glu Lys Phe Asp Pro Asp Ala Thr Gln Phe Met Glu Phe Glu Lys Leu Ser Gln Phe Ala Ala Ala Leu Glu Pro Pro Leu Asn Leu Pro Gln Pro Asn Lys Leu Gln Leu Ile Ala Met Asp Leu Pro Met Val Ser Gly Asp Arg Ile His Cys Leu Asp Ile Leu Phe Ala Phe Thr Lys Arg Val Leu Gly Glu Ser Gly Glu Met Asp Ala Leu Arg Ile Gln Met Glu Glu Arg Phe Met Ala Ser Asn Pro Ser Lys Val Ser Tyr Gln Pro Ile Thr Thr Thr Leu Lys Arg Lys Gln Glu Glu Val Ser Ala Val Ile Ile Gln Page 39

SCN1APCT1.ST25.txt

Arg Ala Tyr Arg Arg His Leu Leu Lys Arg Thr Val Lys Gln Ala 1925 1930 1935

Ser Phe Thr Tyr Asn Lys Asn Lys Ile Lys Gly Gly Ala Asn Leu 1940 1945 1950

Leu Ile Lys Glu Asp Met Ile Ile Asp Arg Ile Asn Glu Asn Ser 1955 1960 1965

Ile Thr Glu Lys Thr Asp Leu Thr Met Ser Thr Ala Ala Cys Pro 1970 1975 1980

Pro Ser Tyr Asp Arg Val Thr Lys Pro Ile Val Glu Lys His Glu 1985 1990 1995

Gln Glu Gly Lys Asp Glu Lys Ala Lys Gly Lys 2000 2005

<210> 5<211> 8381<212> DNA<213> Homo sapiens<400> 5 atactgcaga ggtctctggt gcatgtgtgt atgtgtgcgt ttgtgtgtgt ttgtgtgtct

60

gtgtgttctg ccccagtgag actgcagccc ttgtaaatac tttgacacct tttgcaagaa 120

ggaatctgaa caattgcaac tgaaggcaca ttgttatcat ctcgtctttg ggtgatgctg
180

ttcctcactg cagatggata attttccttt taatcaggaa tttcatatgc agaataaatg 240

gtaattaaaa tgtgcaggat gacaagatgg agcaaacagt gcttgtacca ccaggacctg 300

acagetteaa ettetteace agagaatete ttgeggetat tgaaagaege attgeagaag 360

aaaaggcaaa gaatcccaaa ccagacaaaa aagatgacga cgaaaatggc ccaaagccaa 420

SCN1APCT1.ST25.txt

atagtgactt ggaagctgga aagaaccttc catttattta tggagacatt cctccagaga 480

tggtgtcaga gccctggag gacctggacc cctactatat caataagaaa acttttatag 540

tattgaataa attgaaggcc atcttccggt tcagtgccac ctctgccctg tacattttaa 600

ctcccttcaa tcctcttagg aaaatagcta ttaagatttt ggtacattca ttattcagca 660

tgctaattat gtgcactatt ttgacaaact gtgtgtttat gacaatgagt aaccctcctg
720

attggacaaa gaatgtagaa tacaccttca caggaatata tacttttgaa tcacttataa 780

aaattattgc aaggggattc tgtttagaag attttacttt ccttcgggat ccatggaact 840

ggctcgattt cactgtcatt acatttgcgt acgtcacaga gtttgtggac ctgggcaatg
900

tctcggcatt gagaacattc agagttctcc gagcattgaa gacgatttca gtcattccag
960

gcctgaaaac cattgtggga gccctgatcc agtctgtgaa gaagctctca gatgtaatga 1020

tcctgactgt gttctgtctg agcgtatttg ctctaattgg gctgcagctg ttcatgggca
1080

acctgaggaa taaatgtata caatggcctc ccaccaatgc ttccttggag gaacatagta 1140

tagaaaagaa tataactgtg aattataatg gtacacttat aaatgaaact gtctttgagt 1200

ttgactggaa gtcatatatt caagattcaa gatatcatta tttcctggag ggttttttag
1260

SCN1APCT1.ST25.txt

atgcactact atgtggaaat agctctgatg caggccaatg tccagaggga tatatgtgtg
1320

tgaaagetgg tagaaateee aattatgget acacaagett tgatacette agttgggett 1380

ttttgtcctt gtttcgacta atgactcagg acttctggga aaatctttat caactgacat 1440

tacgtgctgc tgggaaaacg tacatgatat tttttgtatt ggtcattttc ttgggctcat
1500

tctacctaat aaatttgatc ctggctgtgg tggccatggc ctacgaggaa cagaatcagg 1560

ccaccttgga agaagcagaa cagaaagagg ccgaatttca gcagatgatt gaacagctta 1620

aaaagcaaca ggaggcagct cagcaggcag caacggcaac tgcctcagaa cattccagag

agcccagtgc agcaggcagg ctctcagaca gctcatctga agcctctaag ttgagttcca 1740

gggaagagaa agatgaggat gaattccaaa aatctgaatc tgaggacagc atcaggagga 1860

aaggttttcg cttctccatt gaagggaacc gattgacata tgaaaagagg tactcctccc 1920

cacaccagte titgtigage atecgigget cectatitic accaaggega aatageagaa 1980

caageetttt eagetttaga gggegageaa aggatgtggg atetgagaae gaettegeag 2040

atgatgagca cagcacettt gaggataacg agageegtag agatteettg tttgtgeece

SCN1APCT1.ST25.txt

2100

2160

gacgacacgg agagagacgc aacagcaacc tgagtcagac cagtaggtca tcccggatgc

tggcagtgtt tccagcgaat gggaagatgc acagcactgt ggattgcaat ggtgtggttt

2220

ccttggttgg tggaccttca gttcctacat cgcctgttgg acagcttctg ccagaggtga

2280

taatagataa gccagctact gatgacaatg gaacaaccac tgaaactgaa atgagaaaga

2340

gaaggtcaag ttctttccac gtttccatgg actttctaga agatccttcc caaaggcaac

2400

gagcaatgag tatagccagc attctaacaa atacagtaga agaacttgaa gaatccaggc

2460

agaaatgccc accetgttgg tataaatttt ccaacatatt cttaatctgg gactgttctc

2520

catattqqtt aaaaqtqaaa catqttqtca acctqqttqt gatqqaccca tttqttqacc

2580

tggccatcac catctgtatt gtcttaaata ctcttttcat ggccatggag cactatccaa

2640

tgacggacca tttcaataat gtgcttacag taggaaactt ggttttcact gggatcttta

2700

cagcaqaaat gtttctgaaa attattgcca tggatcctta ctattatttc caagaaggct

2760

ggaatatett tgacggtttt attgtgacge ttageetggt agaaettgga etegeeaatg

2820

tggaaggatt atctgttctc cgttcatttc gattgctgcg agttttcaag ttggcaaaat

2880

cttggccaac gttaaatatg ctaataaaga tcatcggcaa ttccgtgggg gctctgggaa

SCN1APCT1_ST25.txt

2940

atttaaccct cgtcttggcc atcatcgtct tcatttttgc cgtggtcggc atgcagctct 3000

ttggtaaaag ctacaaagat tgtgtctgca agatcgccag tgattgtcaa ctcccacgct 3060

ggcacatgaa tgacttette cacteettee tgattgtgtt eegegtgetg tgtggggagt 3120

ggatagagac catgtgggac tgtatggagg ttgctggtca agccatgtgc cttactgtct 3180

tcatgatggt catggtgatt ggaaacctag tggtcctgaa tctctttctg gccttgcttc 3240

tgageteatt tagtgeagae aacettgeag eeactgatga tgataatgaa atgaataate 3300

tccaaattgc tgtggatagg atgcacaaag gagtagctta tgtgaaaaga aaaatatatg 3360

aatttattca acagtccttc attaggaaac aaaagatttt agatgaaatt aaaccacttg 3420

atgatctaaa caacaagaaa gacagttgta tgtccaatca tacaacagaa attgggaaag 3480

atcttgacta tcttaaagat gtaaatggaa ctacaagtgg tataggaact ggcagcagtg 3540

ttgaaaaata cattattgat gaaagtgatt acatgtcatt cataaacaac cccagtctta 3600

ctgtgactgt accaattgct gtaggagaat ctgactttga aaatttaaac acggaagact 3660

ttagtagtga atcggatctg gaagaaagca aagagaaact gaatgaaagc agtagctcat 3720

SCN1APCT1.ST25.txt

cagaaggtag cactgtggac atcggcgcac ctgtagaaga acagcccgta gtggaacctg 3780

aagaaactct tgaaccagaa gcttgtttca ctgaaggctg tgtacaaaga ttcaagtgtt 3840

gtcaaatcaa tgtggaagaa ggcagaggaa aacaatggtg gaacctgaga aggacgtgtt 3900

tccgaatagt tgaacataac tggtttgaga ccttcattgt tttcatgatt ctccttagta 3960

gtggtgctct ggcatttgaa gatatatata ttgatcagcg aaagacgatt aagacgatgt 4020

tggaatatgc tgacaaggtt ttcacttaca ttttcattct ggaaatgctt ctaaaatggg 4080

tggcatatgg ctatcaaaca tatttcacca atgcctggtg ttggctggac ttcttaattg
4140

ttgatgtttc attggtcagt ttaacagcaa atgccttggg ttactcagaa cttggagcca 4200

tcaaatctct caggacacta agagctctga gacctctaag agccttatct cgatttgaag 4260

ggatgagggt ggttgtgaat gcccttttag gagcaattcc atccatcatg aatgtgcttc 4320

tggtttgtct tatattctgg ctaattttca gcatcatggg cgtaaatttg tttgctggca 4380

aattotacca otgtattaac accacaactg gtgacaggtt tgacatogaa gacgtgaata
4440

atcatactga ttgcctaaaa ctaatagaaa gaaatgagac tgctcgatgg aaaaatgtga 4500

aagtaaactt tgataatgta ggatttgggt atctctcttt gcttcaagtt gccacattca 4560

SCN1APCT1.ST25.txt

aaggatggat ggatataatg tatgcagcag ttgattccag aaatgtggaa ctccagccta 4620

agtatgaaaa aagtctgtac atgtatcttt actttgttat tttcatcatc tttgggtcct 4680

tcttcacctt gaacctgttt attggtgtca tcatagataa tttcaaccag cagaaaaaga 4740

agtttggagg tcaagacatc tttatgacag aagaacagaa gaaatactat aatgcaatga 4800

aaaaattagg atcgaaaaaa ccgcaaaagc ctatacctcg accaggaaac aaatttcaag
4860

gaatggtctt tgacttcgta accagacaag tttttgacat aagcatcatg attctcatct 4920

gtcttaacat ggtcacaatg atggtggaaa cagatgacca gagtgaatat gtgactacca 4980

ttttgtcacg catcaatctg gtgttcattg tgctatttac tggagagtgt gtactgaaac 5040

tcatctctct acgccattat tattttacca ttggatggaa tatttttgat tttgtggttg 5100

tcattctctc cattgtaggt atgtttcttg ccgagctgat agaaaagtat ttcgtgtccc 5160

ctaccetgtt ecgagtgate egtettgeta ggattggeeg aateetaegt etgateaaag 5220

gagcaaaggg gatgcgcacg ctgctctttg ctttgatgat gtcccttcct gcgttgttta
5280

acateggeet ectaetette etagteatgt teatetaege eatetttggg atgteeaaet 5340

ttgcctatgt taagagggaa gttgggatcg atgacatgtt caactttgag acctttggca

SCN1APCT1.ST25.txt

5400

acagcatgat ctgcctattc caaattacaa cctctgctgg ctgggatgga ttgctagcac 5460

ccattctcaa cagtaagcca cccgactgtg accctaataa agttaaccct ggaagctcag

5520

ttaagggaga ctgtgggaac ccatctgttg gaattttctt ttttgtcagt tacatcatca

5580

tatccttcct ggttgtggtg aacatgtaca tcgcggtcat cctggagaac ttcagtgttg

5640

ctactgaaga aagtgcagag cctctgagtg aggatgactt tgagatgttc tatgaggttt

5700

gggagaagtt tgatcccgat gcaactcagt tcatggaatt tgaaaaatta tctcagtttg

5760

cagetgeget tgaacegeet eteaatetge cacaaceaaa caaactecag eteattgeca

5820

tqqatttqcc catqqtqagt ggtgaccgga tccactgtct tgatatctta tttgctttta

5880

caaagcqggt tctaggagag agtggagaga tggatgctct acgaatacag atggaagagc

5940

gattcatggc ttccaatcct tccaaggtct cctatcagcc aatcactact actttaaaac

6000

gaaaacaaga gqaagtatct gctgtcatta ttcagcgtgc ttacagacgc caccttttaa

6060

agcgaactgt aaaacaagct tcctttacgt acaataaaaa caaaatcaaa ggtggggcta

6120

atcttcttat aaaaqaaqac atgataattq acaqaataaa tgaaaactct attacagaaa

6180

aaactgatct gaccatgtcc actgcagctt gtccaccttc ctatgaccgg gtgacaaagc

Page 47

SCN1APCT1.ST25.txt

6240

caattgtgga aaaacatgag caagaaggca aagatgaaaa agccaaaggg aaataaatga 6300

aaataaataa aaataattgg gtgacaaatt gtttacagcc tgtgaaggtg atgtattttt 6360

atcaacagga ctcctttagg aggtcaatgc caaactgact gtttttacac aaatctcctt 6420

aaggtcagtg cctacaataa gacagtgacc ccttgtcagc aaactgtgac tctgtgtaaa 6480

ggggagatga cettgacagg aggttactgt teteactace agetgacaet getgaagata 6540

agatgcacaa tggctagtca gactgtaggg accagtttca aggggtgcaa acctgtgatt

ttggggttgt ttaacatgaa acactttagt gtagtaattg tatccactgt ttgcatttca 6660

actgccacat ttgtcacatt tttatggaat ctgttagtgg attcatcttt ttgttaatcc 6720

atgtgtttat tatatgtgac tatttttgta aacgaagttt ctgttgagaa ataggctaag 6780

gacctctata acaggtatgc cacctggggg gtatggcaac cacatggccc tcccagctac .

acaaagtcgt ggtttgcatg agggcatgct gcacttagag atcatgcatg agaaaaagtc 6900

acaagaaaaa caaattotta aatttoacoa tatttotggg aggggtaatt gggtgataag 6960

tggaggtgct ttgttgatct tgttttgcga aatccagccc ctagaccaag tagattattt
7020

SCN1APCT1.ST25.txt

gtgggtaggc cagtaaatct tagcaggtgc aaacttcatt caaatgtttg gagtcataaa 7080

tgttatgttt ctttttgttg tattaaaaaa aaaacctgaa tagtgaatat tgcccctcac

7140

cctccaccgc cagaagactg aattgaccaa aattactctt tataaatttc tgctttttcc

7200

tgcactttgt ttagccatct ttgggctctc agcaaggttg acactgtata tgttaatgaa

7260

atgctattta ttatgtaaat agtcatttta ccctgtggtg cacgtttgag caaacaaata

7320

atgacctaag cacagtattt attgcatcaa atatgtacca caagaaatgt agagtgcaag

7380

ctttacacag gtaataaaat gtattctgta ccatttatag atagtttgga tgctatcaat

7440

gcatgtttat attaccatgc tgctgtatct ggtttctctc actgctcaga atctcattta

7500

tgagaaacca tatgtcagtg gtaaagtcaa ggaaattgtt caacagatct catttattta

7560

agtcattaag caatagtttg cagcacttta acagcttttt ggttattttt acattttaag

7620

tggataacat atggtatata gccagactgt acagacatgt ttaaaaaaaac acactgctta

7680

acctattaaa tatgtgttta gaattttata agcaaatata aatactgtaa aaagtcactt

7740

tattttattt ttcagcatta tgtacataaa tatgaagagg aaattatctt caggttgata

7800

tcacaatcac ttttcttact ttctgtccat agtacttttt catgaaagaa atttgctaaa

7860

SCN1APCT1.ST25.txt

taagacatga aaacaagact gggtagttgt agatttctgc tttttaaatt acatttgcta 7920

attttagatt atttcacaat tttaaggagc aaaataggtt cacgattcat atccaaatta 7980

tgctttgcaa ttggaaaagg gtttaaaatt ttatttatat ttctggtagt acctgtacta 8040

actgaattga aggtagtgct tatgttattt ttgttctttt tttctgactt cggtttatgt 8100

tttcatttct ttggagtaat gctgctctag attgttctaa atagaatgtg ggcttcataa 8160

ttttttttc cacaaaaaca gagtagtcaa cttatatagt caattacatc aggacatttt 8220

gtgtttctta cagaagcaaa ccataggctc ctcttttcct taaaactact tagataaact

gtattcgtga actgcatgct ggaaaatgct actattatgc taaataatgc taaccaacat 8340

ttaaaatgtg caaaactaat aaagattaca ttttttattt t
8381

Phe Thr Arg Glu Ser Leu Ala Ala Ile Glu Arg Arg Ile Ala Glu Glu 20 25 30

Lys Ala Lys Asn Pro Lys Pro Asp Lys Lys Asp Asp Asp Glu Asn Gly 35 40 45

Pro Lys Pro Asn Ser Asp Leu Glu Ala Gly Lys Asn Leu Pro Phe Ile 50 55 60

Page 50

SCN1APCT1.ST25.txt

Tyr Gly Asp Ile Pro Pro Glu Met Val Ser Glu Pro Leu Glu Asp Leu Asp Pro Tyr Tyr Ile Asn Lys Lys Thr Phe Ile Val Leu Asn Lys Leu Lys Ala Ile Phe Arg Phe Ser Ala Thr Ser Ala Leu Tyr Ile Leu Thr Pro Phe Asn Pro Leu Arg Lys Ile Ala Ile Lys Ile Leu Val His Ser Leu Phe Ser Met Leu Ile Met Cys Thr Ile Leu Thr Asn Cys Val Phe Met Thr Met Ser Asn Pro Pro Asp Trp Thr Lys Asn Val Glu Tyr Thr Phe Thr Gly Ile Tyr Thr Phe Glu Ser Leu Ile Lys Ile Ile Ala Arg Gly Phe Cys Leu Glu Asp Phe Thr Phe Leu Arg Asp Pro Trp Asn Trp Leu Asp Phe Thr Val Ile Thr Phe Ala Tyr Val Thr Glu Phe Val Asp Leu Gly Asn Val Ser Ala Leu Arg Thr Phe Arg Val Leu Arg Ala Leu Lys Thr Ile Ser Val Ile Pro Gly Leu Lys Thr Ile Val Gly Ala Leu Ile Gln Ser Val Lys Lys Leu Ser Asp Val Met Ile Leu Thr Val Phe Cys Leu Ser Val Phe Ala Leu Ile Gly Leu Gln Leu Phe Met Gly Asn Leu Arg Asn Lys Cys Ile Gln Trp Pro Pro Thr Asn Ala Ser Leu Glu

Page 51

SCN1APCT1.ST25.txt

Glu His Se 290	r Ile Glu	Lys Asn 295		Thr	Val	Asn	Tyr 300	Asn	Gly	Thr	Leu
Ile Asn Gl 305	u Thr Val	Phe Glu 310	Phe	Asp	Trp	Lys 315	Ser	Tyr	Ile	Gln	Asp 320
Ser Arg Ty	r His Tyr 325		Glu		Phe 330	Leu	Asp	Ala	Leu	Leu 335	Cys
Gly Asn Se	er Ser Asp 340	Ala Gly		Cys 345	Pro	Glu	Gly	Tyr	Met 350	Cys	Val
Lys Ala Gl		Pro Asn	Tyr 360	Gly	Tyr	Thr	Ser	Phe 365	Asp	Thr	Phe
Ser Trp Al	a Phe Leu	Ser Leu 375	Phe	Arg	Leu	Met	Thr 380	Gln	Asp	Phe	Trp
Glu Asn Le 385	eu Tyr Gln	Leu Thr 390	Leu	Arg	Ala	Ala 3 95	Gly	Lys	Thr	Tyr	Met 400
Ile Phe Ph	e Val Leu 405		Phe	Leu	Gly 410	Ser	Phe	Tyr	Leu	Ile 415	Asn
Leu Ile Le	eu Ala Val 420	Val Ala		Ala 425	Tyr	Glu	Glu	Gln	Asn 430	Gln	Ala
Thr Leu Gl		Glu Gln	Lys 440	Glu	Ala	Glu	Phe	Gln 445	Gln	Met	Ile
Glu Gln Le 450	eu Lys Lys	Gln Gln 455	Glu	Ala	Ala	Gln	Gln 460	Ala	Ala	Thr	Ala
Thr Ala Se	er Glu His	Ser Arg 470	Glu	Pro	Ser	Ala 475	Ala	Gly	Arg	Leu	Ser 480
Asp Ser Se	er Ser Glu 485	Ala Ser	Lys	Leu	Ser 490	Ser	Lys	Ser	Ala	Lys 495	Glu
Arg Arg As	n Arg Arg	Lys Lys		Lys age S		Lys	Glu	Gln	Ser	Gly	Gly

510

SCN1APCT1.ST25.txt 500 505

			200					202					510		
Glu	Glu	Lys 515	Asp	Glu	Asp	Glu	Phe 520	Gln	Lys	Ser	Glu	Ser 525	Glu	Asp	Ser
Ile	Arg 530	Arg	Lys	Gly	Phe	Arg 535	Phe	Ser	Ile	Glu	Gly 540	Asn	Arg	Leu	Thr
Tyr 545	Glu	Lys	Arg	Tyr	Ser 550	Ser	Pro	His	Gln	Ser 555	Leu	Leu	Ser	Ile	Arg 560
Gly	Ser	Leu	Phe	Ser 565	Pro	Arg	Arg	Asn	Ser 570	Arg	Thr	Ser	Leu	Phe 575	Ser
Phe	Arg	Gly	Arg 580	Ala	Lys	Asp	Val	Gly 585	Ser	Glu	Asn	Asp	Phe 590	Ala	Asp
Asp	Glu	His 595	Ser	Thr	Phe	Glu	Asp 600	Asn	Glu	Ser	Arg	Arg 605	Asp	Ser	Leu
Phe	Val 610	Pro	Arg	Arg	His	Gly 615	Glu	Arg	Arg	Asn	Ser 620	Asn	Leu	Ser	Gln
Thr 625	Ser	Arg	Ser	Ser	Arg 630	Met	Leu	Ala	Val	Phe 635	Pro	Ala	Asn	Gly	Lys 640
Met	His	Ser	Thr	Val 645	Asp	Cys	Asn	Gly	Val 650	Val	Ser	Leu	Val	Gly 655	Gly
Pro	Ser	Val	Pro 660	Thr	Ser	Pro	Val	Gly 665	Gln	Leu	Leu	Pro	Glu 670	Val	Ile
Ile	Asp	Lys 675	Pro	Ala	Thr	Asp	Asp 680	Asn	Gly	Thr	Thr	Thr 685	Glu	Thr	Glu
Met	Arg 690	Lys	Arg	Arg	Ser	Ser 695	Ser	Phe	His	Val	Ser 700	Met	Asp	Phe	Leu
Glu 705	Asp	Pro	Ser	Gln	Arg 710	Gln	Arg	Ala	Met	Ser 715	Ile	Ala	Ser	Ile	Leu 720

SCN1APCT1.ST25.txt

Thr Asn Thr Val Glu Glu Leu Glu Glu Ser Arg Gln Lys Cys Pro Pro 725 730 735

Cys Trp Tyr Lys Phe Ser Asn Ile Phe Leu Ile Trp Asp Cys Ser Pro 740 745 750

Tyr Trp Leu Lys Val Lys His Val Val Asn Leu Val Val Met Asp Pro 755 760 765

Phe Val Asp Leu Ala Ile Thr Ile Cys Ile Val Leu Asn Thr Leu Phe 770 775 780

Met Ala Met Glu His Tyr Pro Met Thr Asp His Phe Asn Asn Val Leu 785 790 795 800

Thr Val Gly Asn Leu Val Phe Thr Gly Ile Phe Thr Ala Glu Met Phe 805 810 815

Leu Lys Ile Ile Ala Met Asp Pro Tyr Tyr Tyr Phe Gln Glu Gly Trp 820 825 830

Asn Ile Phe Asp Gly Phe Ile Val Thr Leu Ser Leu Val Glu Leu Gly 835 840 845

Leu Ala Asn Val Glu Gly Leu Ser Val Leu Arg Ser Phe Arg Leu Leu 850 855 860

Arg Val Phe Lys Leu Ala Lys Ser Trp Pro Thr Leu Asn Met Leu Ile 865 870 875 880

Lys Ile Ile Gly Asn Ser Val Gly Ala Leu Gly Asn Leu Thr Leu Val 885 890 895

Leu Ala Ile Ile Val Phe Ile Phe Ala Val Val Gly Met Gln Leu Phe 900 905 910

Gly Lys Ser Tyr Lys Asp Cys Val Cys Lys Ile Ala Ser Asp Cys Gln 915 920 925

Leu Pro Arg Trp His Met Asn Asp Phe Phe His Ser Phe Leu Ile Val 930 935 940

SCN1APCT1.ST25.txt

Phe Arg Val Leu Cys Gly Glu Trp Ile Glu Thr Met Trp Asp Cys Met 945 950 955 960

Glu Val Ala Gly Gln Ala Met Cys Leu Thr Val Phe Met Met Val Met 965 970 975

Val Ile Gly Asn Leu Val Val Leu Asn Leu Phe Leu Ala Leu Leu Leu 980 985 990

Ser Ser Phe Ser Ala Asp Asn Leu Ala Ala Thr Asp Asp Asp Asn Glu 995 1000 1005

Met Asn Asn Leu Gln Ile Ala Val Asp Arg Met His Lys Gly Val 1010 1015 1020

Ala Tyr Val Lys Arg Lys Ile Tyr Glu Phe Ile Gln Gln Ser Phe 1025 1030 1035

Ile Arg Lys Gln Lys Ile Leu Asp Glu Ile Lys Pro Leu Asp Asp 1040 1045 1050

Leu Asn Asn Lys Lys Asp Ser Cys Met Ser Asn His Thr Thr Glu 1055 1060 1065

Ile Gly Lys Asp Leu Asp Tyr Leu Lys Asp Val Asn Gly Thr Thr 1070 1080

Ser Gly Ile Gly Thr Gly Ser Ser Val Glu Lys Tyr Ile Ile Asp 1085 1090 1095

Glu Ser Asp Tyr Met Ser Phe Ile Asn Asn Pro Ser Leu Thr Val 1100 1105 1110

Thr Val Pro Ile Ala Val Gly Glu Ser Asp Phe Glu Asn Leu Asn 1115 1120 1125

Thr Glu Asp Phe Ser Ser Glu Ser Asp Leu Glu Glu Ser Lys Glu 1130 1135 1140

Lys Leu Asn Glu Ser Ser Ser Ser Glu Gly Ser Thr Val Asp 1145 1150 1155

Page 55

SCN1APCT1.ST25.txt

Ile	Gly 1160		Pro	Val	Glu	Glu 1165	Gln	Pro	Val	Val	Glu 1170	Pro	Glu	Glu
Thr	Leu 1175	Glu	Pro	Glu	Ala	Cys 1180		Thr	Glu	Gly	Cys 1185	Val	Gln	Arg
Phe	Lys 1190	_	Cys	Gln	Ile	Asn 1195		Glu	Glu	Gly	Arg 1200	Gly	Lys	Gln
Trp	Trp 1205		Leu	Arg	Arg	Thr 1210	Cys	Phe	Arg	Ile	Val 1215	Glu	His	Asn
Trp	Phe 1220	Glu	Thr	Phe	Ile	Val 1225	Phe	Met	Ile	Leu	Leu 1230	Ser	Ser	Gly
Ala	Leu 1235		Phe	Glu	Asp	Ile 1240	Tyr	Ile	Asp	Gln	Arg 1245	Lys	Thr	Ile
Lys	Thr 1250	Met	Leu	Glu	Tyr	Ala 1255	Asp	Lys	Val	Phe	Thr 1260	Tyr	Ile	Phe
Ile	Leu 1265		Met	Leu	Leu	Lys 1270		Val	Ala	Tyr	Gly 1275	Tyr	Gln	Thr
Tyr	Phe 1280	Thr	Asn	Ala	Trp	Cys 1285	Trp	Leu	Asp	Phe	Leu 1290	Ile	Val	Asp
Val	Ser 1295	Leu	Val	Ser	Leu	Thr 1300	Ala	Asn	Ala	Leu	Gly 1305	Tyr	Ser	Glu
Leu	Gly 1310	Ala	Ile	Lys	Ser	Leu 1315	Arg	Thr	Leu	Arg	Ala 1320	Leu	Arg	Pro
Leu	Arg 1325	Ala	Leu	Ser	Arg	Phe 1330	Glu	Gly	Met	Arg	Val 1335	Val	Val	Asn
Ala	Leu 1340	Leu	Gly	Ala	Ile	Pro 1345	Ser	Ile	Met	Asn	Val 1350	Leu	Leu	Val
Cys	Leu .	Ile	Phe	Trp	Leu	Ile		Ser e 56	Ile	Met	Gly	Val	Asn	Leu

1365

SCN1APCT1.ST25.txt 1355 1360

	1333					1300					1303			
Phe	Ala 1370		Lys	Phe		His 1375		Ile	Asn	Thr	Thr 1380	Thr	Gly	Asp
Arg	Phe 1385	Asp	Ile	Glu	Asp	Val 1390	Asn	Asn	His	Thr	Asp 1395	Cys	Leu	Lys
Leu	Ile 1400	Glu	Arg	Asn	Glu	Thr 1405	Ala	Arg	Trp	Lys	Asn 1410	Val	Lys	Val
Asn	Phe 1415		Asn	Val	Gly	Phe 1420		Tyr	Leu	Ser	Leu 1425	Leu	Gln	Val
Ala	Thr 1430	Phe	Lys	Gly	Trp	Met 1435		Ile	Met	Tyr	Ala 1440	Ala	Val	Asp
Ser	Arg 1445	Asn	Val	Glu	Leu	Gln 1450		Lys	Tyr	Glu	Lys 1455	Ser	Leu	Tyr
Met	Tyr 1460	Leu	Tyr	Phe	Val	Ile 1465	Phe	Ile	Ile	Phe	Gly 1470	Ser	Phe	Phe
Thr	Leu 1475	Asn	Leu	Phe	Ile	Gly 1480	Val	Ile	Ile	Asp	Asn 1485	Phe	Asn	Gln
Gln	Lys 1490		Lys	Phe		Gly 1495	Gln	Asp	Ile	Phe	Met 1500	Thr	Glu	Glu
Gln	Lys 1505	Lys	Tyr	Tyr	Asn	Ala 1510	Met	Lys	Lys	Leu	Gly 1515	Ser	Lys	Lys
Pro	Gln 1520	Lys	Pro	Ile	Pro	Arg 1525	Pro	Gly	Asn	Lys	Phe 1530	Gln	Gly	Met
Val	Phe 1535	Asp	Phe	Val	Thr	Arg 1540	Gln	Val	Phe	Asp	Ile 1545	Ser	Ile	Met
Ile	Leu 1550	Ile	Cys	Leu	Asn	Met 1555	Val	Thr	Met	Met	Val 1560	Glu	Thr	Asp

	SCN1APCT1.ST25.txt													
Asp	Gln 1565		Glu	Tyr	Val	Thr 1570	Thr				Arg 1575		Asn	Leu
Val	Phe 1580	Ile	Val	Leu	Phe	Thr 1585	Gly	Glu	Cys	Val	Leu 1590	Lys	Leu	Ile
Ser	Leu 1595	Arg	His	Tyr	Tyr	Phe 1600	Thr	Ile	Gly	Trp	Asn 1605	I1e	Phe	Asp
Phe	Val 1610		Val	Ile	Leu	Ser 1615	I1e	Val	G1y	Met	Phe 1620	Leu	Ala	Glu
Leu	Ile 1625	Glu	Lys	Tyr	Phe	Val 1630	Ser	Pro	Thr	Leu	Phe 1635	Arg	Val	Ile
Arg	Leu 1640		Arg	Ile	Gly	Arg 1645		Leu	Arg	Leu	Ile 1650	Lys	Gly	Ala
Lys	Gly 1655	Met	Arg	Thr	Leu	Leu 1660	Phe	Ala	Leu	Met	Met 1665	Ser	Leu	Pro
Ala	Leu 1670	Phe	Asn	Ile	Gly	Leu 1675	Leu	Leu	Phe	Leu	Val 1680	Met	Phe	Ile
Tyr	Ala 1685	Ile	Phe	Gly	Met	Ser 1690	Asn	Phe	Ala	Tyr	Val 1695	Lys	Arg	Glu
Val	Gly 1700		Asp	Asp	Met	Phe 1705	Asn	Phe	Glu	Thr	Phe 1710	Gly	Asn	Ser
Met	Ile 1715	Cys	Leu	Phe	Gln	Ile 1720	Thr	Thr	Ser	Ala	Gly 1725	Trp	Asp	Gly
Leu	Leu 1730	Ala	Pro	Ile	Leu	Asn 1735	Ser	Lys	Pro	Pro	Asp 1740	Cys	Asp	Pro
Asn	Lys 1745	Val	Asn	Pro	Gly	Ser 1750	Ser	Val	Lys	Gly	Asp 1755	Cys	Gly	Asn
Pro	Ser 1760	Val	Gly	Ile	Phe	Phe 1765	Phe	Val	Ser	Tyr	Ile 1770	Ile	Ile	Ser

SCN1APCT1.ST25.txt

Phe	Leu 1775		Val	Val	Asn	Met 1780	Tyr	Ile	Ala	Val	Ile 1785	Leu	Glu	Asn
Phe	Ser 1790	Val	Ala	Thr	Glu	Glu 1795		Ala	Glu	Pro	Leu 1800	Ser	Glu	Asp
Asp	Phe 1805	Glu	Met	Phe	Tyr	Glu 1810	Val	Trp	Glu	Lys	Phe 1815	Asp	Pro	Asp
Ala	Thr 1820	Gln	Phe	Met	Glu	Phe 1825	Glu	Lys	Leu	Ser	Gln 1830	Phe	Ala	Ala
Ala	Leu 1835		Pro	Pro	Leu	Asn 1840	Leu	Pro	Gln	Pro	Asn 1845	Lys	Leu	Gln
Leu	Ile 1850	Ala	Met	Asp	Leu	Pro 1855	Met	Val	Ser	Gly	Asp 1860	Arg	Ile	His
Cys	Leu 1865		Ile	Leu	Phe	Ala 1870	Phe	Thr	Lys	Arg	Val 1875	Leu	Gly	Glu
Ser	Gly 1880	Glu	Met	Asp	Ala	Leu 1885	Arg	Ile	Gln	Met	Glu 1890	Glu	Arg	Phe
Met	Ala 1895	Ser	Asn	Pro	Ser	Lys 1900	Val	Ser	Tyr	Gln	Pro 1905	Ile	Thr	Thr
Thr	Leu 1910	Lys	Arg	Lys	Gln	Glu 1915	Glu	Val	Ser	Ala	Val 1920	Ile	Ile	Gln
Arg	Ala 1925	Tyr	Arg	Arg	His	Leu 1930	Leu	Lys	Arg	Thr	Val 1935	Lys	Gln	Ala
Ser	Phe 1940	Thr	Tyr	Asn	Lys	Asn 1945	Lys	Ile	Lys	Gly	Gly 1950	Ala	Asn	Leu
Leu	Ile 1955	Lys	Glu	Asp	Met	Ile 1960	Ile	Asp	Arg	Ile	Asn 1965	Glu	Asn	Ser
Ile	Thr 1970	Glu	Lys	Thr	Asp	Leu 1975		Met e 59	Ser	Thr	Ala 1980	Ala	Cys	Pro
							- ~9							

SCN1APCT1.ST25.txt

Pro Ser Tyr Asp Arg Val Thr Lys Pro Ile Val Glu Lys His Glu 1985 1990 1995

Gln Glu Gly Lys Asp Glu Lys Ala Lys Gly Lys 2000 2005

<210> 7<211> 8381<212> DNA<213> Homo sapiens<400> 7 atactgcaga ggtctctggt gcatgtgtgt atgtgtgcgt ttgtgtgtgt ttgtgtgtct

60

gtgtgttctg ccccagtgag actgcagccc ttgtaaatac tttgacacct tttgcaagaa
120

ggaatctgaa caattgcaac tgaaggcaca ttgttatcat ctcgtctttg ggtgatgctg
180

ttcctcactg cagatggata attttccttt taatcaggaa tttcatatgc agaataaatg 240

gtaattaaaa tgtgcaggat gacaagatgg agcaaacagt gcttgtacca ccaggacctg

acagetteaa ettetteace agagaatete ttgeggetat tgaaagaege attgeagaag 360

aaaaggcaaa gaatcccaaa ccagacaaaa aagatgacga cgaaaatggc ccaaagccaa 420

atagtgactt ggaagctgga aagaaccttc catttattta tggagacatt cctccagaga 480

tggtgtcaga gccctggag gacctggacc cctactatat caataagaaa acttttatag

tattgaataa attgaaggcc atcttccggt tcagtgccac ctctgccctg tacattttaa 600

ctcccttcaa tcctcttagg aaaatagcta ttaagatttt ggtacattca ttattcagca 660

SCN1APCT1.ST25.txt

tgctaattat gtgcactatt ttgacaaact gtgtgtttat gacaatgagt aacceteetg
720

attggacaaa gaatgtagaa tacacettca caggaatata tacttttgaa tcacttataa 780

aaattattgc aaggggattc tgtttagaag attttacttt ccttcgggat ccatggaact

ggctcgattt cactgtcatt acatttgcgt acgtcacaga gtttgtggac ctgggcaatg 900

tctcggcatt gagaacattc agagttctcc gagcattgaa gacgatttca gtcattccag 960

gcctgaaaac cattgtggga gccctgatcc agtctgtgaa gaagctctca gatgtaatga

tcctgactgt gttctgtctg agcgtatttg ctctaattgg gctgcagctg ttcatgggca 1080

acctgaggaa taaatgtata caatggcctc ccaccaatgc ttccttggag gaacatagta 1140

tagaaaagaa tataactgtg aattataatg gtacacttat aaatgaaact gtctttgagt 1200

ttgactggaa gtcatatatt caagattcaa gatatcatta tttcctggag ggttttttag
1260

atgcactact atgtggaaat agctctgatg caggccaatg tccagaggga tatatgtgtg
1320

tgaaagctgg tagaaatccc aattatggct acacaagctt tgataccttc agttgggctt 1380

ttttgtcctt gtttcgacta atgactcagg acttctggga aaatctttat caactgacat 1440

tacgtgctgc tgggaaaacg tacatgatat tttttgtatt ggtcattttc ttgggctcat 1500

SCN1APCT1.ST25.txt

tctacctaat aaatttgatc ctggctgtgg tggccatggc ctacgaggaa cagaatcagg

ccaccttgga agaagcagaa cagaaagagg ccgaatttca gcagatgatt gaacagctta
1620

aaaagcaaca ggaggcagct cagcaggcag caacggcaac tgcctcagaa cattccagag 1680

agcccagtgc agcaggcagg ctctcagaca gctcatctga agcctctaag ttgagttcca
1740

gggaagagaa agatgaggat gaattccaaa aatctgaatc tgaggacagc atcaggagga 1860

aaggttttcg cttctccatt gaagggaacc gattgacata tgaaaagagg tactcctccc 1920

cacaccagtc tttgttgagc atccgtggct ccctattttc accaaggcga aatagcagaa 1980

caagcetttt cagetttaga gggegageaa aggatgtggg atetgagaac gaettegeag 2040

atgatgagca cagcaccttt gaggataacg agagccgtag agattccttg tttgtgcccc 2100

gacgacacgg agagagacgc aacagcaacc tgagtcagac cagtaggtca tcccggatgc 2160

tggcagtgtt tccagcgaat gggaagatgc acagcactgt ggattgcaat ggtgtggttt 2220

cettggttgg tggacettca gtteetacat egeetgttgg acagettetg ecagaggtga 2280

taatagataa gccagctact gatgacaatg gaacaaccac tgaaactgaa atgagaaaga

SCN1APCT1.ST25.txt

2340

2460

gaaggtcaag ttctttccac gtttccatgg actttctaga agatccttcc caaaggcaac 2400

gagcaatgag tatagccagc attctaacaa atacagtaga agaacttgaa gaatccaggc

agaaatgccc accetgttgg tataaatttt ccaacatatt cttaatctgg gactgttctc 2520

catattggtt aaaagtgaaa catgttgtca acctggttgt gatggaccca tttgttgacc 2580

tggccatcac catctgtatt gtcttaaata ctcttttcat ggccatggag cactatccaa 2640

tgacggacca tttcaataat gtgcttacag taggaaactt ggttttcact gggatcttta
2700

cagcagaaat gtttctgaaa attattgcca tggatcctta ctattatttc caagaaggct
2760

ggaatatett tgaeggtttt attgtgagge ttageetggt agaaettgga etegeeaatg 2820

tggaaggatt atctgttctc cgttcatttc gattgctgcg agttttcaag ttggcaaaat 2880

cttggccaac gttaaatatg ctaataaaga tcatcggcaa ttccgtgggg gctctgggaa 2940

atttaaccct cgtcttggcc atcatcgtct tcatttttgc cgtggtcggc atgcagctct 3000

ttggtaaaag ctacaaagat tgtgtctgca agatcgccag tgattgtcaa ctcccacgct 3060

ggcacatgaa tgacttcttc cactccttcc tgattgtgtt ccgcgtgctg tgtggggagt 3120

ggatagagac catgtgggac tgtatggagg ttgctggtca agccatgtgc cttactgtct

Page 63

SCN1APCT1.ST25.txt

3180

tcatgatggt catggtgatt ggaaacctag tggtcctgaa tctctttctg gccttgcttc 3240

tgagctcatt tagtgcagac aaccttgcag ccactgatga tgataatgaa atgaataatc 3300

tccaaattgc tgtggatagg atgcacaaag gagtagctta tgtgaaaaga aaaatatatg 3360

aatttattca acagtccttc attaggaaac aaaagatttt agatgaaatt aaaccacttg 3420

atgatctaaa caacaagaaa gacagttgta tgtccaatca tacaacagaa attgggaaag 3480

atcttgacta tcttaaagat gtaaatggaa ctacaagtgg tataggaact ggcagcagtg . 3540

ttgaaaaata cattattgat gaaagtgatt acatgtcatt cataaacaac cccagtctta 3600

ctgtgactgt accaattgct gtaggagaat ctgactttga aaatttaaac acggaagact 3660

ttagtagtga atcggatctg gaagaaagca aagagaaact gaatgaaagc agtagctcat 3720

cagaaggtag cactgtggac atcggcgcac ctgtagaaga acagcccgta gtggaacctg

aagaaactct tgaaccagaa gcttgtttca ctgaaggctg tgtacaaaga ttcaagtgtt 3840

gtcaaatcaa tgtggaagaa ggcagaggaa aacaatggtg gaacctgaga aggacgtgtt 3900

tccgaatagt tgaacataac tggtttgaga ccttcattgt tttcatgatt ctccttagta 3960

SCN1APCT1.ST25.txt

gtggtgctct ggcatttgaa gatatatata ttgatcagcg aaagacgatt aagacgatgt 4020

tggaatatgc tgacaaggtt ttcacttaca ttttcattct ggaaatgctt ctaaaatggg 4080

tggcatatgg ctatcaaaca tatttcacca atgcctggtg ttggctggac ttcttaattg 4140

ttgatgtttc attggtcagt ttaacagcaa atgccttggg ttactcagaa cttggagcca 4200

tcaaatctct caggacacta agagctctga gacctctaag agccttatct cgatttgaag 4260

ggatgaggt ggttgtgaat gcccttttag gagcaattcc atccatcatg aatgtgcttc 4320

tggtttgtct tatattctgg ctaattttca gcatcatggg cgtaaatttg tttgctggca 4380

aattctacca ctgtattaac accacaactg gtgacaggtt tgacatcgaa gacgtgaata 4440

atcatactga ttgcctaaaa ctaatagaaa gaaatgagac tgctcgatgg aaaaatgtga 4500

aagtaaactt tgataatgta ggatttgggt atctctcttt gcttcaagtt gccacattca 4560

aaggatggat ggatataatg tatgcagcag ttgattccag aaatgtggaa ctccagccta 4620

agtatgaaaa aagtctgtac atgtatcttt actttgttat tttcatcatc tttgggtcct 4680

tcttcacctt gaacctgttt attggtgtca tcatagataa tttcaaccag cagaaaaaga 4740

agtttggagg tcaagacatc tttatgacag aagaacagaa gaaatactat aatgcaatga 4800

SCN1APCT1.ST25.txt

aaaaattagg atcgaaaaaa ccgcaaaagc ctatacctcg accaggaaac aaatttcaag
4860

gaatggtett tgaettegta accagacaag tttttgaeat aagcateatg atteteatet 4920

gtcttaacat ggtcacaatg atggtggaaa cagatgacca gagtgaatat gtgactacca 4980

ttttgtcacg catcaatctg gtgttcattg tgctatttac tggagagtgt gtactgaaac 5040

tcatctctct acgccattat tattttacca ttggatggaa tatttttgat tttgtggttg 5100

tcattctctc cattgtaggt atgtttcttg ccgagctgat agaaaagtat ttcgtgtccc 5160

ctaccetgtt ecgagtgate egtettgeta ggattggeeg aateetaegt etgateaaag 5220

gagcaaaggg gatccgcacg ctgctctttg ctttgatgat gtcccttcct gcgttgttta
5280

acateggeet cetaetette etagteatgt teatetaege eatetttggg atgteeaact 5340

ttgcctatgt taagagggaa gttgggatcg atgacatgtt caactttgag acctttggca 5400

acagcatgat ctgcctattc caaattacaa cctctgctgg ctgggatgga ttgctagcac 5460

ccatteteaa cagtaageea eeegaetgtg accetaataa agttaaeeet ggaageteag 5520

ttaagggaga ctgtgggaac ccatctgttg gaattttctt ttttgtcagt tacatcatca 5580

tatecttect ggttgtggtg aacatgtaca tegeggteat cetggagaac tteagtgttg

SCN1APCT1.ST25.txt

5640

ctactgaaga aagtgcagag cctctgagtg aggatgactt tgagatgttc tatgaggttt 5700

gggagaagtt tgatcccgat gcaactcagt tcatggaatt tgaaaaatta tctcagtttg 5760

cagctgcgct tgaaccgcct ctcaatctgc cacaaccaaa caaactccag ctcattgcca 5820

tggatttgcc catggtgagt ggtgaccgga tccactgtct tgatatctta tttgctttta
5880

caaagcgggt tctaggagag agtggagaga tggatgctct acgaatacag atggaagagc 5940

gattcatggc ttccaatcct tccaaggtct cctatcagcc aatcactact actttaaaac 6000

gaaaacaaga ggaagtatct gctgtcatta ttcagcgtgc ttacagacgc caccttttaa 6060

agcgaactgt aaaacaagct tcctttacgt acaataaaaa caaaatcaaa ggtggggcta 6120

atcttcttat aaaagaagac atgataattg acagaataaa tgaaaactct attacagaaa 6180

aaactgatct gaccatgtcc actgcagctt gtccaccttc ctatgaccgg gtgacaaagc 6240

caattgtgga aaaacatgag caagaaggca aagatgaaaa agccaaaggg aaataaatga 6300

aaataaataa aaataattgg gtgacaaatt gtttacagcc tgtgaaggtg atgtattttt 6360

atcaacagga ctcctttagg aggtcaatgc caaactgact gtttttacac aaatctcctt 6420

aaggtcagtg cctacaataa gacagtgacc ccttgtcagc aaactgtgac tctgtgtaaa

Page 67

SCN1APCT1.ST25.txt

6480

ggggagatga cettgacagg aggttactgt tetcactace agetgacaet getgaagata 6540

agatgcacaa tggctagtca gactgtaggg accagtttca aggggtgcaa acctgtgatt 6600

ttggggttgt ttaacatgaa acactttagt gtagtaattg tatccactgt ttgcatttca 6660

actgccacat ttgtcacatt tttatggaat ctgttagtgg attcatcttt ttgttaatcc 6720

atgtgtttat tatatgtgac tatttttgta aacgaagttt ctgttgagaa ataggctaag 6780

gacctctata acaggtatgc cacctggggg gtatggcaac cacatggccc tcccagctac 6840

acaaagtcgt ggtttgcatg agggcatgct gcacttagag atcatgcatg agaaaaagtc 6900

acaagaaaaa caaattotta aatttoacca tatttotggg aggggtaatt gggtgataag 6960

tggaggtgct ttgttgatct tgttttgcga aatccagccc ctagaccaag tagattattt 7020

gtgggtaggc cagtaaatct tagcaggtgc aaacttcatt caaatgtttg gagtcataaa 7080

tgttatgttt ctttttgttg tattaaaaaa aaaacctgaa tagtgaatat tgcccctcac 7140

cctccaccgc cagaagactg aattgaccaa aattactctt tataaatttc tgctttttcc 7200

tgcactttgt ttagccatct ttgggctctc agcaaggttg acactgtata tgttaatgaa 7260

SCN1APCT1.ST25.txt

atgctattta ttatgtaaat agtcatttta ccctgtggtg cacgtttgag caaacaaata 7320

atgacctaag cacagtattt attgcatcaa atatgtacca caagaaatgt agagtgcaag

ctttacacag gtaataaaat gtattctgta ccatttatag atagtttgga tgctatcaat 7440

gcatgtttat attaccatgc tgctgtatct ggtttctctc actgctcaga atctcattta 7500

tgagaaacca tatgtcagtg gtaaagtcaa ggaaattgtt caacagatct catttattta
7560

agtcattaag caatagtttg cagcacttta acagcttttt ggttattttt acattttaag 7620

tggataacat atggtatata gccagactgt acagacatgt ttaaaaaaac acactgctta
7680

acctattaaa tatgtgttta gaattttata agcaaatata aatactgtaa aaagtcactt 7740

tattttattt ttcagcatta tgtacataaa tatgaagagg aaattatctt caggttgata 7800

tcacaatcac ttttcttact ttctgtccat agtacttttt catgaaagaa atttgctaaa 7860

taagacatga aaacaagact gggtagttgt agatttctgc tttttaaatt acatttgcta
7920

attttagatt atttcacaat tttaaggagc aaaataggtt cacgattcat atccaaatta 7980

tgctttgcaa ttggaaaagg gtttaaaatt ttatttatat ttctggtagt acctgtacta . 8040

actgaattga aggtagtgct tatgttattt ttgttctttt tttctgactt cggtttatgt 8100

SCN1APCT1.ST25.txt

tttcatttct ttggagtaat gctgctctag attgttctaa atagaatgtg ggcttcataa 8160

ttttttttc cacaaaaca gagtagtcaa cttatatagt caattacatc aggacatttt 8220

gtgtttctta cagaagcaaa ccataggctc ctcttttcct taaaactact tagataaact 8280

gtattcgtga actgcatgct ggaaaatgct actattatgc taaataatgc taaccaacat 8340

ttaaaatgtg caaaactaat aaagattaca ttttttattt t 8381

<210> 8<211> 8381<212> DNA<213> Homo sapiens<400> 8 atactgcaga ggtctctggt gcatgtgtgt atgtgtgcgt ttgtgtgtgt ttgtgtgtct 60

gtgtgttctg ccccagtgag actgcagccc ttgtaaatac tttgacacct tttgcaagaa 120

ggaatctgaa caattgcaac tgaaggcaca ttgttatcat ctcgtctttg ggtgatgctg
180

ttcctcactg cagatggata attttccttt taatcaggaa tttcatatgc agaataaatg 240

gtaattaaaa tgtgcaggat gacaagatgg agcaaacagt gcttgtacca ccaggacctg

acagetteaa ettetteace agagaatete ttgeggetat tgaaagaege attgeagaag 360

aaaaggcaaa gaatcccaaa ccagacaaaa aagatgacga cgaaaatggc ccaaagccaa 420

atagtgactt ggaagctgga aagaaccttc catttattta tggagacatt cctccagaga 480

SCN1APCT1.ST25.txt

tggtgtcaga gccctggag gacctggacc cctactatat caataagaaa acttttatag 540

tattgaataa attgaaggcc atcttccggt tcagtgccac ctctgccctg tacattttaa 600

ctcccttcaa tcctcttagg aaaatagcta ttaagatttt ggtacattca ttattcagca 660

tgctaattat gtgcactatt ttgacaaact gtgtgtttat gacaatgagt aaccctcctg
720

attggacaaa gaatgtagaa tacaccttca caggaatata tacttttgaa tcacttataa 780

aaattattgc aaggggattc tgtttagaag attttacttt ccttcgggat ccatggaact 840

ggctcgattt cactgtcatt acatttgcgt acgtcacaga gtttgtggac ctgggcaatg

tctcggcatt gagaacattc agagttctcc gagcattgaa gacgatttca gtcattccag
960

gcctgaaaac cattgtggga gccctgatcc agtctgtgaa gaagctctca gatgtaatga 1020

tcctgactgt gttctgtctg agcgtatttg ctctaattgg gctgcagctg ttcatgggca
1080

acctgaggaa taaatgtata caatggcctc ccaccaatgc ttccttggag gaacatagta 1140

tagaaaagaa tataactgtg aattataatg gtacacttat aaatgaaact gtctttgagt 1200

ttgactggaa gtcatatatt caagattcaa gatatcatta tttcctggag ggttttttag
1260

atgcactact atgtggaaat agctctgatg caggccaatg tccagaggga tatatgtgtg

SCN1APCT1.ST25.txt

tgaaagctgg tagaaatccc aattatggct acacaagctt tgataccttc agttgggctt 1380

ttttgtcctt gtttcgacta atgactcagg acttctggga aaatctttat caactgacat 1440

tacgtgctgc tgggaaaacg tacatgatat tttttgtatt ggtcattttc ttgggctcat 1500

tctacctaat aaatttgatc ctggctgtgg tggccatggc ctacgaggaa cagaatcagg
1560

ccaccttgga agaagcagaa cagaaagagg ccgaatttca gcagatgatt gaacagctta 1620

aaaagcaaca ggaggcagct cagcaggcag caacggcaac tgcctcagaa cattccagag
1680

ageceagtge ageaggeagg eteteagaea geteatetga ageetetaag ttgagtteea
1740

gggaagagaa agatgaggat gaattccaaa aatctgaatc tgaggacagc atcaggagga 1860

aaggttttcg cttctccatt gaagggaacc ga**tt**gacata tgaaaagagg tactcctccc 1920

cacaccagtc tttgttgagc atccgtggct ccctattttc accaaggcga aatagcagaa 1980

caageetttt cagetttaga gggegageaa aggatgtggg atetgagaac gaettegeag 2040

atgatgagca cagcaccttt gaggataacg agagccgtag agattccttg tttgtgcccc 2100

gacgacacgg agagagacgc aacagcaacc tgagtcagac cagtaggtca tcccggatgc

SCN1APCT1.ST25.txt

2160

tggcagtgtt tccagcgaat gggaagatgc acagcactgt ggattgcaat ggtgtggttt

2220

ccttggttgg tggaccttca gttcctacat cgcctgttgg acagcttctg ccagaggtga

2280

taatagataa gccagctact gatgacaatg gaacaaccac tgaaactgaa atgagaaaga

2340

gaaggtcaag ttctttccac gtttccatgg actttctaga agatccttcc caaaggcaac

2400

gagcaatgag tatagccagc attctaacaa atacagtaga agaacttgaa gaatccaggc

2460

agaaatgccc accetgttgg tataaatttt ccaacatatt cttaatctgg gactgttctc

2520

catattggtt aaaagtgaaa catgttgtca acctggttgt gatggaccca tttgttgacc

2580

tggccatcac catctgtatt gtcttaaata ctcttttcat ggccatggag cactatccaa

2640

tgacggacca tttcaataat gtgcttacag taggaaactt ggttttcact gggatcttta

2700

cagcagaaat gtttctgaaa attattgcca tggatcctta ctattatttc caagaaggct

2760

ggaatatett tgaeggtttt attgtgaege ttageetggt agaacttgga etegeeaatg

2820

tggaaggatt atctgttctc cgttcatttc gattgctgcg agttttcaag ttggcaaaat

2880

cttggccaac gttaaatatg ctaataaaga tcatcggcaa ttccgtgggg gctctgggaa

2940

atttaaccct cgtcttggcc atcatcgtct tcatttttgc cgtggtcggc atgcagctct

SCN1APCT1.ST25.txt

3000

ttggtaaaag ctacaaagat tgtgtctgca agatcgccag tgattgtcaa ctcccacgct

3060

ggcacatgaa tgacttcttc cactccttcc tgattgtgtt ccgcgtgctg tgtggggagt

3120

ggatagagac catgtgggac tgtatggagg ttgccggtca agccatgtgc cttactgtct

3180

tcatgatggt catggtgatt ggaaacctag tggtcctgaa tctctttctg gccttgcttc

3240

tgagctcatt tagtgcagac aaccttgcag ccactgatga tgataatgaa atgaataatc

3300

tccaaattgc tgtggatagg atgcacaaag gagtagctta tgtgaaaaga aaaatatatg

3360

aatttattca acagtccttc attaggaaac aaaagatttt agatgaaatt aaaccacttg

3420

atgatctaaa caacaagaaa gacagttgta tgtccaatca tacaacagaa attgggaaag

3480

atcttgacta tcttaaagat gtaaatggaa ctacaagtgg tataggaact ggcagcagtg

3540

ttgaaaaata cattattgat gaaagtgatt acatgtcatt cataaacaac cccagtctta

3600

ctgtgactgt accaattgct gtaggagaat ctgactttga aaatttaaac acggaagact

3660

ttagtagtga atcggatctg gaagaaagca aagagaaact gaatgaaagc agtagctcat

3720

cagaaggtag cactgtggac atcggcgcac ctgtagaaga acagcccgta gtggaacctg

3780

SCN1APCT1.ST25.txt

aagaaactct tgaaccagaa gcttgtttca ctgaaggctg tgtacaaaga ttcaagtgtt 3840

gtcaaatcaa tgtggaagaa ggcagaggaa aacaatggtg gaacctgaga aggacgtgtt 3900

tccgaatagt tgaacataac tggtttgaga ccttcattgt tttcatgatt ctccttagta 3960

gtggtgctct ggcatttgaa gatatatata ttgatcagcg aaagacgatt aagacgatgt 4020

tggaatatgc tgacaaggtt ttcacttaca ttttcattct ggaaatgctt ctaaaatggg 4080

tggcatatgg ctatcaaaca tatttcacca atgcctggtg ttggctggac ttcttaattg 4140

ttgatgtttc attggtcagt ttaacagcaa atgccttggg ttactcagaa cttggagcca 4200

tcaaatctct caggacacta agagctctga gacctctaag agccttatct cgatttgaag 4260

ggatgagggt ggttgtgaat gcccttttag gagcaattcc atccatcatg aatgtgcttc 4320

tggtttgtct tatattctgg ctaattttca gcatcatggg cgtaaatttg tttgctggca 4380

aattctacca ctgtattaac accacaactg gtgacaggtt tgacatcgaa gacgtgaata 4440

atcatactga ttgcctaaaa ctaatagaaa gaaatgagac tgctcgatgg aaaaatgtga 4500

aagtaaactt tgataatgta ggatttgggt atctctcttt gcttcaagtt gccacattca 4560

aaggatggat ggatataatg tatgcagcag ttgattccag aaatgtggaa ctccagccta 4620

SCN1APCT1.ST25.txt

agtatgaaaa aagtetgtac atgtatettt aetttgttat ttteateate tttgggteet 4680

tcttcacctt gaacctgttt attggtgtca tcatagataa tttcaaccag cagaaaaaga 4740

agtttggagg tcaagacatc tttatgacag aagaacagaa gaaatactat aatgcaatga 4800

aaaaattagg atcgaaaaaa ccgcaaaagc ctatacctcg accaggaaac aaatttcaag 4860

gaatggtctt tgacttcgta accagacaag tttttgacat aagcatcatg attctcatct 4920

gtcttaacat ggtcacaatg atggtggaaa cagatgacca gagtgaatat gtgactacca 4980

ttttgtcacg catcaatctg gtgttcattg tgctatttac tggagagtgt gtactgaaac 5040

tcatctctct acgccattat tattttacca ttggatggaa tatttttgat tttgtggttg 5100

tcattctctc cattgtaggt atgtttcttg ccgagctgat agaaaagtat ttcgtgtccc 5160

ctaccetgtt cegagtgate egtettgeta ggattggeeg aateetaegt etgateaaag 5220

gagcaaaggg gatccgcacg ctgctctttg ctttgatgat gtcccttcct gcgttgttta 5280

acateggeet cetaetette etagteatgt teatetaege eatetttggg atgteeaact 5340

ttgcctatgt taagagggaa gttgggatcg atgacatgtt caactttgag acctttggca 5400

acagcatgat ctgcctattc caaattacaa cctctgctgg ctgggatgga ttgctagcac

SCN1APCT1.ST25.txt

5460

ccattctcaa cagtaagcca cccgactgtg accctaataa agttaaccct ggaagctcag

5520

ttaagggaga ctgtgggaac ccatctgttg gaattttctt ttttgtcagt tacatcatca

5580

tatccttcct ggttgtggtg aacatgtaca tcgcggtcat cctggagaac ttcagtgttg

5640

ctactgaaga aagtgcagag cctctgagtg aggatgactt tgagatgttc tatgaggttt

5700

gggagaagtt tgatcccgat gcaactcagt tcatggaatt tgaaaaatta tctcagttttg

5760

cagctgcgct tgaaccgcct ctcaatctgc cacaaccaaa caaactccag ctcattgcca

5820

tggatttgcc catggtgagt ggtgaccgga tccactgtct tgatatctta tttgctttta

5880

caaagcgggt tctaggagag agtggagaga tggatgctct acgaatacag atggaagagc

5940

gattcatggc ttccaatcct tccaaggtct cctatcagcc aatcactact actttaaaac

6000

gaaaacaaga ggaagtatct gctgtcatta ttcagcgtgc ttacagacgc caccttttaa

6060

agcgaactgt aaaacaagct tcctttacgt acaataaaaa caaaatcaaa ggtggggcta

6120

atcttcttat aaaagaagac atgataattg acagaataaa tgaaaactct attacagaaa

6180

aaactgatct gaccatgtcc actgcagctt gtccaccttc ctatgaccgg gtgacaaagc

6240

caattgtgga aaaacatgag caagaaggca aagatgaaaa agccaaaggg aaataaatga

Page 77

SCN1APCT1.ST25.txt

6300

aaataaataa aaataattgg gtgacaaatt gtttacagcc tgtgaaggtg atgtattttt 6360

atcaacagga ctcctttagg aggtcaatgc caaactgact gtttttacac aaatctcctt 6420

aaggtcagtg cctacaataa gacagtgacc ccttgtcagc aaactgtgac tctgtgtaaa 6480

ggggagatga cettgacagg aggttactgt tetcactace agetgacaet getgaagata 6540

agatgcacaa tggctagtca gactgtaggg accagtttca aggggtgcaa acctgtgatt 6600

ttggggttgt ttaacatgaa acactttagt gtagtaattg tatccactgt ttgcatttca 6660

actgccacat ttgtcacatt tttatggaat ctgttagtgg attcatcttt ttgttaatcc 6720

atgtgtttat tatatgtgac tatttttgta aacgaagttt ctgttgagaa ataggctaag 6780

gacctctata acaggtatgc cacctggggg gtatggcaac cacatggccc tcccagctac 6840

acaaagtcgt ggtttgcatg agggcatgct gcacttagag atcatgcatg agaaaaagtc 6900

acaagaaaaa caaattotta aatttoacoa tatttotggg aggggtaatt gggtgataag 6960

tggaggtgct ttgttgatct tgttttgcga aatccagccc ctagaccaag tagattattt 7020

gtgggtaggc cagtaaatct tagcaggtgc aaacttcatt caaatgtttg gagtcataaa . 7080

SCN1APCT1.ST25.txt

tgttatgttt ctttttgttg tattaaaaaa aaaacctgaa tagtgaatat tgcccctcac 7140

cctccaccgc cagaagactg aattgaccaa aattactctt tataaatttc tgctttttcc 7200

tgcactttgt ttagccatct ttgggctctc agcaaggttg acactgtata tgttaatgaa 7260

atgctattta ttatgtaaat agtcatttta ccctgtggtg cacgtttgag caaacaaata 7320

atgacctaag cacagtattt attgcatcaa atatgtacca caagaaatgt agagtgcaag 7380

ctttacacag gtaataaaat gtattctgta ccatttatag atagtttgga tgctatcaat 7440

gcatgtttat attaccatgc tgctgtatct ggtttctctc actgctcaga atctcattta 7500

tgagaaacca tatgtcagtg gtaaagtcaa ggaaattgtt caacagatct catttattta 7560

agtcattaag caatagtttg cagcacttta acagcttttt ggttattttt acattttaag 7620

tggataacat atggtatata gccagactgt acagacatgt ttaaaaaaac acactgctta 7680

acctattaaa tatgtgttta gaattttata agcaaatata aatactgtaa aaagtcactt 7740

tattttattt ttcagcatta tgtacataaa tatgaagagg aaattatctt caggttgata 7800

tcacaatcac ttttcttact ttctgtccat agtacttttt catgaaagaa atttgctaaa 7860

taagacatga aaacaagact gggtagttgt agatttctgc tttttaaatt acatttgcta
7920

SCN1APCT1.ST25.txt

attttagatt atttcacaat tttaaggagc aaaataggtt cacgattcat atccaaatta 7980

tgctttgcaa ttggaaaagg gtttaaaatt ttatttatat ttctggtagt acctgtacta 8040

actgaattga aggtagtgct tatgttattt ttgttctttt tttctgactt cggtttatgt 8100

tttcatttct ttggagtaat gctgctctag attgttctaa atagaatgtg ggcttcataa 8160

ttttttttc cacaaaaaca gagtagtcaa cttatatagt caattacatc aggacatttt
8220

gtgtttctta cagaagcaaa ccataggctc ctcttttcct taaaactact tagataaact

gtattcgtga actgcatgct ggaaaatgct actattatgc taaataatgc taaccaacat 8340

ttaaaatgtg caaaactaat aaagattaca ttttttattt t 8381

<210> 9<211> 8381<212> DNA<213> Homo sapiens<400> 9 atactgcaga ggtctctggt gcatgtgtgt atgtgtgcgt ttgtgtgtgt ttgtgtgtct 60

gtgtgttctg ccccagtgag actgcagccc ttgtaaatac tttgacacct tttgcaagaa 120

ggaatctgaa caattgcaac tgaaggcaca ttgttatcat ctcgtctttg ggtgatgctg
180

ttcctcactg cagatggata attttccttt taatcaggaa tttcatatgc agaataaatg 240

gtaattaaaa tgtgcaggat gacaagatgg agcaaacagt gcttgtacca ccaggacctg

SCN1APCT1.ST25.txt

acagetteaa ettetteace agagaatete ttgeggetat tgaaagaege attgeagaag 360

aaaaggcaaa gaatcccaaa ccagacaaaa aagatgacga cgaaaatggc ccaaagccaa

420

atagtgactt ggaagctgga aagaaccttc catttattta tggagacatt cctccagaga

480

tggtgtcaga gccctggag gacctggacc cctactatat caataagaaa acttttatag

540

tattgaataa attgaaggcc atcttccggt tcagtgccac ctctgccctg tacattttaa

600

ctcccttcaa tcctcttagg aaaatagcta ttaagatttt ggtacattca ttattcagca

660

tgctaattat gtgcactatt ttgacaaact gtgtgtttat gacaatgagt aacceteetg

720

attggacaaa gaatgtagaa tacaccttca caggaatata tacttttgaa tcacttataa

780

aaattattgc aaggggattc tgtttagaag attttacttt ccttcgggat ccatggaact

840

ggctcgattt cactgtcatt acatttgcgt acgtcacaga gtttgtggac ctgggcaatg

900

totoggcatt gagaacattc agagttotoc gagcattgaa gacgatttca gtcattccag

960

gcctgaaaac cattgtggga gccctgatcc agtctgtgaa gaagctctca gatgtaatga

1020

tcctgactgt gttctgtctg agcgtatttg ctctaattgg gctgcagctg ttcatgggca

1080

acctgaggaa taaatgtata caatggcctc ccaccaatgc ttccttggag gaacatagta

1140

SCN1APCT1.ST25.txt

tagaaaagaa tataactgtg aattataatg gtacacttat aaatgaaact gtctttgagt 1200

ttgactggaa gtcatatatt caagattcaa gatatcatta tttcctggag ggttttttag
1260

atgcactact atgtggaaat agctctgatg caggccaatg tccagaggga tatatgtgtg
1320

tgaaagetgg tagaaateee aattatgget acacaagett tgatacette agttgggett 1380

ttttgtcctt gtttcgacta atgactcagg acttctggga aaatctttat caactgacat 1440

tacgtgctgc tgggaaaacg tacatgatat tttttgtatt ggtcattttc ttgggctcat
1500

tctacctaat aaatttgatc ctggctgtgg tggccatggc ctacgaggaa cagaatcagg

ccaccttgga agaagcagaa cagaaagagg ccgaatttca gcagatgatt gaacagctta 1620

aaaagcaaca **gg**aggcagct cagcaggcag caacggcaac tgcctcagaa cattccagag 1680

agcccagtgc agcaggcagg ctctcagaca gctcatctga agcctctaag ttgagttcca

gggaagagaa agatgaggat gaattccaaa aatctgaatc tgaggacagc atcaggagga 1860

aaggttttcg cttctccatt gaagggaacc gattgacata tgaaaagagg tactcctccc 1920

cacaccaqte tttgttgage atceqtgget ceetatttte accaaggega aatageagaa

SCN1APCT1.ST25.txt

1980

caageetttt cagetttaga gggegageaa aggatgtggg atetgagaae gaettegeag

2040

atgatgagca cagcaccttt gaggataacg agagccgtag agattccttg tttgtgcccc

2100

gacgacacgg agagagacgc aacagcaacc tgagtcagac cagtaggtca tcccggatgc

2160

tggcagtgtt tccagcgaat gggaagatgc acagcactgt ggattgcaat ggtgtggttt

2220

ccttggttgg tggaccttca gttcctacat cgcctgttgg acagcttctg ccagaggtga

2280

taatagataa gccagctact gatgacaatg gaacaaccac tgaaactgaa atgagaaaga

2340

gaaggtcaag ttctttccac gtttccatgg actttctaga agatccttcc caaaggcaac

2400

qaqcaatqaq tataqccagc attctaacaa atacagtaga agaacttgaa gaatccaggc

2460

agaaatgccc accctgttgg tataaatttt ccaacatatt cttaatctgg gactgttctc

2520

catattggtt aaaagtgaaa catgttgtca acctggttgt gatggaccca tttgttgacc

2580

tggccatcac catctgtatt gtcttaaata ctcttttcat ggccatggag cactatccaa

2640

tgacggacca tttcaataat gtgcttacag taggaaactt ggttttcact gggatcttta

2700

cagcagaaat gtttctgaaa attattgcca tggatcctta ctattatttc caagaaggct

2760

ggaatatett tgacggtttt attgtgacge ttageetggt agaacttgga etegeeaatg

SCN1APCT1.ST25.txt

2820

tggaaggatt atctgttctc cgttcatttc gattgctgcg agttttcaag ttggcaaaat 2880

cttggccaac gttaaatatg ctaataaaga tcatcggcaa ttccgtgggg gctctgggaa 2940

atttaaccct cgtcttggcc atcatcgtct tcatttttgc cgtggtcggc atgcagctct 3000

ttggtaaaag ctacaaagat tgtgtctgca agatcgccag tgattgtcaa ctcccacgct 3060

ggcacatgaa tgacttcttc cactccttcc tgattgtgtt ccgcgtgctg tgtggggagt 3120

ggatagagac catgtgggac tgtatggagg ttgctggtca agccatgtgc cttactgtct 3180

tcatgatggt catggtgatt ggaaacctag tggtcctgaa tctctttctg gccttgcttc 3240

tgagctcatt tagtgcagac aaccttgcag ccactgatga tgataatgaa atgaataatc 3300

tccaaattgc tgtggatagg atgcacaaag gagtagctta tgtgaaaaga aaaatatatg 3360

aatttattca acagtccttc attaggaaac aaaagatttt agatgaaatt aaaccacttg 3420

atgatctaaa caacaagaaa gacagttgta tgtccaatca tacagcagaa attgggaaag 3480

atcttgacta tcttaaagat gtaaatggaa ctacaagtgg tataggaact ggcagcagtg 3540

ttgaaaaata cattattgat gaaagtgatt acatgtcatt cataaacaac cccagtctta 3600

SCN1APCT1.ST25.txt

ctgtgactgt accaattgct gtaggagaat ctgactttga aaatttaaac acggaagact 3660

ttagtagtga atcggatctg gaagaaagca aagagaaact gaatgaaagc agtagctcat 3720

cagaaggtag cactgtggac atcggcgcac ctgtagaaga acagcccgta gtggaacctg 3780

aagaaactct tgaaccagaa gcttgtttca ctgaaggctg tgtacaaaga ttcaagtgtt 3840

gtcaaatcaa tgtggaagaa ggcagaggaa aacaatggtg gaacctgaga aggacgtgtt 3900

tccgaatagt tgaacataac tggtttgaga ccttcattgt tttcatgatt ctccttagta 3960

gtggtgctct ggcatttgaa gatatatata ttgatcagcg aaagacgatt aagacgatgt 4020

tggaatatgc tgacaaggtt ttcacttaca ttttcattct ggaaatgctt ctaaaatggg 4080

tggcatatgg ctatcaaaca tatttcacca atgcctggtg ttggctggac ttcttaattg 4140

ttgatgtttc attggtcagt ttaacagcaa atgccttggg ttactcagaa cttggagcca 4200

tcaaatctct caggacacta agagctctga gacctctaag agccttatct cgatttgaag 4260

ggatgagggt ggttgtgaat gcccttttag gagcaattcc atccatcatg aatgtgcttc 4320

tggtttgtct tatattctgg ctaattttca gcatcatggg cgtaaatttg tttgctggca

aattctacca ctgtattaac accacaactg gtgacaggtt tgacatcgaa gacgtgaata 4440

SCN1APCT1.ST25.txt

atcatactga ttgcctaaaa ctaatagaaa gaaatgagac tgctcgatgg aaaaatgtga 4500

aagtaaactt tgataatgta ggatttgggt atctctcttt gcttcaagtt gccacattca 4560

aaggatggat ggatataatg tatgcagcag ttgattccag aaatgtggaa ctccagccta 4620

agtatgaaaa aagtotgtac atgtatottt actttgttat tttcatcatc tttgggtoot 4680

tcttcacctt gaacctgttt attggtgtca tcatagataa tttcaaccag cagaaaaaga 4740

agtttggagg tcaagacatc tttatgacag aagaacagaa gaaatactat aatgcaatga 4800

aaaaattagg atcgaaaaaa ccgcaaaagc ctatacctcg accaggaaac aaatttcaag
4860

gaatggtctt tgacttcgta accagacaag tttttgacat aagcatcatg attctcatct 4920

gtcttaacat ggtcacaatg atggtggaaa cagatgacca gagtgaatat gtgactacca 4980

ttttgtcacg catcaatctg gtgttcattg tgctatttac tggagagtgt gtactgaaac 5040

tcatctctct acgccattat tattttacca ttggatggaa tatttttgat tttgtggttg 5100

tcattctctc cattgtaggt atgtttcttg ccgagctgat agaaaagtat ttcgtgtccc 5160

ctaccetgtt ecgagtgate egtettgeta ggattggeeg aateetaegt etgateaaag 5220

gagcaaaggg gatccgcacg ctgctctttg ctttgatgat gtcccttcct gcgttgttta

SCN1APCT1.ST25.txt

5280

acateggeet cetaetette etagteatgt teatetaege eatetttggg atgteeaaet 5340

ttgcctatgt taagagggaa gttgggatcg atgacatgtt caactttgag acctttggca 5400

acagcatgat ctgcctattc caaattacaa cctctgctgg ctgggatgga ttgctagcac 5460

ccatteteaa eagtaageea eeegaetgtg accetaataa agttaaeeet ggaageteag 5520

ttaagggaga ctgtgggaac ccatctgttg gaattttctt ttttgtcagt tacatcatca 5580

tatcetteet ggttgtggtg aacatgtaca tegeggteat eetggagaac tteagtgttg 5640

ctactgaaga aagtgcagag cctctgagtg aggatgactt tgagatgttc tatgaggttt 5700

gggagaagtt tgatcccgat gcaactcagt tcatggaatt tgaaaaatta tctcagtttg 5760

cagetgeget tgaacegeet etcaatetge cacaaceaaa caaacteeag etcattgeca 5820

tggatttgcc catggtgagt ggtgaccgga tccactgtct tgatatctta tttgctttta
5880

caaagcgggt tctaggagag agtggagaga tggatgctct acgaatacag atggaagagc 5940

gattcatggc ttccaatcct tccaaggtct cctatcagcc aatcactact actttaaaac 6000

gaaaacaaga ggaagtatct gctgtcatta ttcagcgtgc ttacagacgc caccttttaa 6060

agcgaactgt aaaacaagct tcctttacgt acaataaaaa caaaatcaaa ggtggggcta
Page 87

SCN1APCT1.ST25.txt

6120

atcttcttat aaaagaagac atgataattg acagaataaa tgaaaactct attacagaaa 6180

aaactgatct gaccatgtcc actgcagctt gtccaccttc ctatgaccgg gtgacaaagc 6240

caattgtgga aaaacatgag caagaaggca aagatgaaaa agccaaaggg aaataaatga 6300

aaataaataa aaataattgg gtgacaaatt gtttacagcc tgtgaaggtg atgtattttt 6360

atcaacagga ctcctttagg aggtcaatgc caaactgact gtttttacac aaatctcctt 6420

aaggtcagtg cctacaataa gacagtgacc ccttgtcagc aaactgtgac tctgtgtaaa 6480

ggggagatga ccttgacagg aggttactgt tctcactacc agctgacact gctgaagata 6540

agatgcacaa tggctagtca gactgtaggg accagtttca aggggtgcaa acctgtgatt 6600

ttggggttgt ttaacatgaa acactttagt gtagtaattg tatccactgt ttgcatttca 6660

actgccacat ttgtcacatt tttatggaat ctgttagtgg attcatcttt ttgttaatcc 6720

atgtgtttat tatatgtgac tatttttgta aacgaagttt ctgttgagaa ataggctaag 6780

gacctctata acaggtatgc cacctggggg gtatggcaac cacatggccc tcccagctac 6840

acaaagtcgt ggtttgcatg agggcatgct gcacttagag atcatgcatg agaaaaagtc

SCN1APCT1.ST25.txt

acaagaaaaa caaattotta aatttoacca tatttotggg aggggtaatt gggtgataag 6960

tggaggtgct ttgttgatct tgttttgcga aatccagccc ctagaccaag tagattattt 7020

gtgggtaggc cagtaaatct tagcaggtgc aaacttcatt caaatgtttg gagtcataaa 7080

tgttatgttt ctttttgttg tattaaaaaa aaaacctgaa tagtgaatat tgcccctcac 7140

cctccaccgc cagaagactg aattgaccaa aattactctt tataaatttc tgctttttcc 7200

tgcactttgt ttagccatct ttgggctctc agcaaggttg acactgtata tgttaatgaa 7260

atgctattta ttatgtaaat agtcatttta ccctgtggtg cacgtttgag caaacaaata 7320

atgacctaag cacagtattt attgcatcaa atatgtacca caagaaatgt agagtgcaag 7380

ctttacacag gtaataaaat gtattctgta ccatttatag atagtttgga tgctatcaat 7440

gcatgtttat attaccatgc tgctgtatct ggtttctctc actgctcaga atctcattta 7500

tgagaaacca tatgtcagtg gtaaagtcaa ggaaattgtt caacagatct catttattta 7560

agtcattaag caatagtttg cagcacttta acagcttttt ggttattttt acattttaag 7620

tggataacat atggtatata gccagactgt acagacatgt ttaaaaaaac acactgctta 7680

acctattaaa tatgtgttta gaattttata agcaaatata aatactgtaa aaagtcactt 7740

SCN1APCT1.ST25.txt

tattttattt ttcagcatta tgtacataaa tatgaagagg aaattatctt caggttgata 7800

tcacaatcac ttttcttact ttctgtccat agtacttttt catgaaagaa atttgctaaa 7860

taagacatga aaacaagact gggtagttgt agatttctgc tttttaaatt acatttgcta
7920

attttagatt atttcacaat tttaaggagc aaaataggtt cacgattcat atccaaatta 7980

tgctttgcaa ttggaaaagg gtttaaaatt ttatttatat ttctggtagt acctgtacta 8040

actgaattga aggtagtgct tatgttattt ttgttctttt tttctgactt cggtttatgt 8100

tttcatttct ttggagtaat gctgctctag attgttctaa atagaatgtg ggcttcataa 8160

ttttttttc cacaaaaca gagtagtcaa cttatatagt caattacatc aggacatttt 8220

gtgtttctta cagaagcaaa ccataggctc ctcttttcct taaaactact tagataaact 8280

gtattcgtga actgcatgct ggaaaatgct actattatgc taaataatgc taaccaacat 8340

ttaaaatgtg caaaactaat aaagattaca ttttttattt t 8381

Phe Thr Arg Glu Ser Leu Ala Ala Ile Glu Arg Arg Ile Ala Glu Glu 20 25 30

SCN1APCT1.ST25.txt

Lys Ala Lys Asn Pro Lys Pro Asp Lys Lys Asp Asp Asp Glu Asn Gly 35 40 45

Pro Lys Pro Asn Ser Asp Leu Glu Ala Gly Lys Asn Leu Pro Phe Ile 50 55 60

Tyr Gly Asp Ile Pro Pro Glu Met Val Ser Glu Pro Leu Glu Asp Leu 65 70 75 80

Asp Pro Tyr Tyr Ile Asn Lys Lys Thr Phe Ile Val Leu Asn Lys Leu 85 90 95

Lys Ala Ile Phe Arg Phe Ser Ala Thr Ser Ala Leu Tyr Ile Leu Thr 100 105 110

Pro Phe Asn Pro Leu Arg Lys Ile Ala Ile Lys Ile Leu Val His Ser 115 120 125

Leu Phe Ser Met Leu Ile Met Cys Thr Ile Leu Thr Asn Cys Val Phe 130 135 140

Met Thr Met Ser Asn Pro Pro Asp Trp Thr Lys Asn Val Glu Tyr Thr 145 150 155 160

Phe Thr Gly Ile Tyr Thr Phe Glu Ser Leu Ile Lys Ile Ile Ala Arg 165 170 175

Gly Phe Cys Leu Glu Asp Phe Thr Phe Leu Arg Asp Pro Trp Asn Trp 180 185 190

Leu Asp Phe Thr Val Ile Thr Phe Ala Tyr Val Thr Glu Phe Val Asp 195 200 205

Leu Gly Asn Val Ser Ala Leu Arg Thr Phe Arg Val Leu Arg Ala Leu 210 215 220

Lys Thr Ile Ser Val Ile Pro Gly Leu Lys Thr Ile Val Gly Ala Leu 225 230 235 240

Ile Gln Ser Val Lys Lys Leu Ser Asp Val Met Ile Leu Thr Val Phe 245 250 255

SCN1APCT1.ST25.txt

Cys	Leu	Ser	Val 260	Phe	Ala	Leu	Ile	Gly 265	Leu	Gln	Leu	Phe	Met 270	Gly	Asn
Leu	Arg	Asn 275	Lys	Cys	Ile	Gln	Trp 280	Pro	Pro	Thr	Asn	Ala 285	Ser	Leu	Glu
Glu	His 290	Ser	Ile	Glu	Lys	Asn 295	Ile	Thr	Val	Asn	Tyr 300	Asn	Gly	Thr	Leu
Ile 305	Asn	Glu	Thr	Val	Phe 310	Glu	Phe	Asp	Trp	Lys 315	Ser	Tyr	Ile	Gln	Asp 320
Ser	Arg	Tyr	His	Tyr 325	Phe	Leu	Glu	Gly	Phe 330	Leu	Asp	Ala	Leu	Leu 335	Cys
Gly	Asn	Ser	Ser 340	Asp	Ala	Gly	Gln	Cys 345	Pro	Glu	Gly	Tyr	Met 350	Cys	Val
Lys	Ala	Gly 355	Arg	Asn	Pro	Asn	Tyr 360	Gly	Tyr	Thr	Ser	Phe 365	Asp	Thr	Phe
Ser	Trp 370	Ala	Phe	Leu	Ser	Leu 375	Phe	Arg	Leu	Met	Thr 380	Gln	Asp	Phe	Trp
Glu 385	Asn	Leu	Tyr	Gln	Leu 390	Thr	Leu	Arg	Ala	Ala 395	Gly	Lys	Thr	Tyr	Met 400
Ile	Phe	Phe	Val	Leu 405	Val	Ile	Phe	Leu	Gly 410	Ser	Phe	Tyr	Leu	Ile 415	Asn
Leu	Ile	Leu	Ala 420	Val	Val	Ala	Met	Ala 425	Tyr	Glu	Glu	Gln	Asn 430	Gln	Ala
Thr	Leu	Glu 435	Glu	Ala	Glu	Gln	Lys 440	Glu	Ala	Glu	Phe	Gln 445	Gln	Met	Ile
Glu	Gln 450	Leu	Lys	Lys	Gln	Gln 455	Glu	Ala	Ala	Gln	Gln 460	Ala	Ala	Thr	Ala
Thr	Ala	Ser	Glu	His	Ser	Arg		Pro age		Ala	Ala	Gly	Arg	Leu	Ser

490

					470	SCN	11APC	T1.5	T25.						400
465	465									475					480
Asp	Ser	Ser	Ser	Glu	Ala	Ser	Lys	Leu	Ser	Ser	Lys	Ser	Ala	Lys	Glu

Arg Arg Asn Arg Arg Lys Lys Arg Lys Gln Lys Glu Gln Ser Gly Gly 500 505 510

485

Glu Glu Lys Asp Glu Asp Glu Phe Gln Lys Ser Glu Ser Glu Asp Ser 515 520 525

Ile Arg Arg Lys Gly Phe Arg Phe Ser Ile Glu Gly Asn Arg Leu Thr 530 540

Tyr Glu Lys Arg Tyr Ser Ser Pro His Gln Ser Leu Leu Ser Ile Arg 545 550 550 560

Gly Ser Leu Phe Ser Pro Arg Arg Asn Ser Arg Thr Ser Leu Phe Ser 565 570 575

Phe Arg Gly Arg Ala Lys Asp Val Gly Ser Glu Asn Asp Phe Ala Asp 580 585 590

Asp Glu His Ser Thr Phe Glu Asp Asn Glu Ser Arg Asp Ser Leu 595 600 605

Phe Val Pro Arg Arg His Gly Glu Arg Arg Asn Ser Asn Leu Ser Gln 610 620

Thr Ser Arg Ser Ser Arg Met Leu Ala Val Phe Pro Ala Asn Gly Lys 625 630 635 640

Met His Ser Thr Val Asp Cys Asn Gly Val Val Ser Leu Val Gly Gly $645 \hspace{1cm} 650 \hspace{1cm} 655$

Pro Ser Val Pro Thr Ser Pro Val Gly Gln Leu Leu Pro Glu Val Ile 660 665 670

Ile Asp Lys Pro Ala Thr Asp Asp Asn Gly Thr Thr Thr Glu Thr Glu 675 680 685

SCN1APCT1.ST25.txt Met Arg Lys Arg Arg Ser Ser Ser Phe His Val Ser Met Asp Phe Leu Glu Asp Pro Ser Gln Arg Gln Arg Ala Met Ser Ile Ala Ser Ile Leu Thr Asn Thr Val Glu Glu Leu Glu Glu Ser Arg Gln Lys Cys Pro Pro Cys Trp Tyr Lys Phe Ser Asn Ile Phe Leu Ile Trp Asp Cys Ser Pro Tyr Trp Leu Lys Val Lys His Val Val Asn Leu Val Val Met Asp Pro Phe Val Asp Leu Ala Ile Thr Ile Cys Ile Val Leu Asn Thr Leu Phe Met Ala Met Glu His Tyr Pro Met Thr Asp His Phe Asn Asn Val Leu Thr Val Gly Asn Leu Val Phe Thr Gly Ile Phe Thr Ala Glu Met Phe Leu Lys Ile Ile Ala Met Asp Pro Tyr Tyr Tyr Phe Gln Glu Gly Trp Asn Ile Phe Asp Gly Phe Ile Val Thr Leu Ser Leu Val Glu Leu Gly Leu Ala Asn Val Glu Gly Leu Ser Val Leu Arg Ser Phe Arg Leu Leu Arg Val Phe Lys Leu Ala Lys Ser Trp Pro Thr Leu Asn Met Leu Ile Lys Ile Ile Gly Asn Ser Val Gly Ala Leu Gly Asn Leu Thr Leu Val Leu Ala Ile Ile Val Phe Ile Phe Ala Val Val Gly Met Gln Leu Phe

SCN1APCT1.ST25.txt

Gly Lys Ser Tyr Lys Asp Cys Val Cys Lys Ile Ala Ser Asp Cys Gln 915 920 925

Leu Pro Arg Trp His Met Asn Asp Phe Phe His Ser Phe Leu Ile Val 930 935 940

Phe Arg Val Leu Cys Gly Glu Trp Ile Glu Thr Met Trp Asp Cys Met 945 950 955 960

Glu Val Ala Gly Gln Ala Met Cys Leu Thr Val Phe Met Met Val Met 965 970 975

Val Ile Gly Asn Leu Val Val Leu Asn Leu Phe Leu Ala Leu Leu 980 985 990

Ser Ser Phe Ser Ala Asp Asn Leu Ala Ala Thr Asp Asp Asp Asn Glu 995 1000 1005

Met Asn Asn Leu Gln Ile Ala Val Asp Arg Met His Lys Gly Val 1010 \$1015\$ 1020

Ala Tyr Val Lys Arg Lys Ile Tyr Glu Phe Ile Gln Gln Ser Phe 1025 1030 1035

Ile Arg Lys Gln Lys Ile Leu Asp Glu Ile Lys Pro Leu Asp Asp 1040 1045 1050

Leu Asn Asn Lys Lys Asp Ser Cys Met Ser Asn His Thr Ala Glu 1055 1060 1065

Ile Gly Lys Asp Leu Asp Tyr Leu Lys Asp Val Asn Gly Thr Thr 1070 1080

Ser Gly Ile Gly Thr Gly Ser Ser Val Glu Lys Tyr Ile Ile Asp 1085 1090 1095

Glu Ser Asp Tyr Met Ser Phe Ile Asn Asn Pro Ser Leu Thr Val 1100 1105 1110

Thr Val Pro Ile Ala Val Gly Glu Ser Asp Phe Glu Asn Leu Asn 1115 1120 1125

SCN1APCT1.ST25.txt

Thr	Glu 1130	Asp	Phe	Ser	Ser	Glu 1135		Asp	Leu	Glu	Glu 1140	Ser	Lys	Glu
Lys	Leu 1145	Asn	Glu	Ser	Ser	Ser 1150		Ser	Glu	Gly	Ser 1155	Thr	Va1	Asp
Ile	Gly 1160	Ala	Pro	Val	Glu	Glu 1165	Gln	Pro	Val	Val	Glu 1170	Pro	G1u	Glu
Thr	Leu 1175	Glu	Pro	Glu	Ala	Cys 1180	Phe	Thr	Glu	Gly	Cys 1185	Va1	Gln	Arg
Phe	Lys 1190	Cys	Cys	Gln	Ile	Asn 1195	Val	Glu	Glu	Gly	Arg 1200	Gly	Lys	Gln
Trp	Trp 1205	Asn	Leu	Arg	Arg	Thr 1210	Cys	Phe	Arg	Ile	Val 1215	Glu	His	Asn
Trp	Phe 1220	Glu	Thr	Phe	Ile	Val 1225	Phe	Met	Ile	Leu	L eu 1230	Ser	Ser	Gly
Ala	Leu 1235	Ala	Phe	Glu	Asp	Ile 1240	Tyr	Ile	Asp	Gln	Arg 1245	Lys	Thr	Ile
Lys	Thr 1250	Met	Leu	Glu	Tyr	Ala 1255	Asp	Lys	Val	Phe	Thr 1260	Tyr	Ile	Phe
Ile	Leu 1265	Glu	Met	Leu	Leu	Lys 1270	Trp	Val	Ala	Tyr	Gly 1275	Tyr	Gln	Thr
Tyr	Phe 1280	Thr	Asn	Ala	Trp	Cys 1285	Trp	Leu	Asp	Phe	Leu 1290	Ile	Val	Asp
Val	Ser 1295	Leu	Val	Ser	Leu	Thr 1300	Ala	Asn	Ala	Leu	Gly 1305	Tyr	Ser	Glu
Leu	Gly 1310	Ala	Ile	Lys	Ser	Leu 1315	Arg	Thr	Leu	Arg	Ala 1320	Leu	Arg	Pro
Leu	Arg	Ala	Leu	Ser	Arg	Phe		Gly e 96	Met	Arg	Val	Va1	Val	Asn

1335

SCN1APCT1.ST25.txt 1325 1330

	± 32 3													
Ala	Leu 1340		Gly	Ala	Ile	Pro 1345		Ile	Met	Asn	Val 1350	Leu	Leu	Val
Cys	Leu 1355	Ile	Phe	Trp	Leu	Ile 1360	Phe	Ser	Ile	Met	Gly 1365	Val	Asn	Leu
Phe	Ala 1370	Gly	Lys	Phe	Tyr	His 1375		Ile	Asn	Thr	Thr 1380	Thr	Gly	Asp
Arg	Phe 1385	Asp	Ile	Glu	Asp	Val 1390	Asn	Asn	His	Thr	Asp 1395	Cys	Leu	Lys
Leu	Ile 1400	Glu	Arg	Asn	Glu	Thr 1405		Arg	Trp	Lys	Asn 1410	Val	Lys	Val
Asn	Phe 1415		Asn	Val	Gly	Phe 1420	Gly	Tyr	Leu	Ser	Leu 1425	Leu	Gln	Val
Ala	Thr 1430	Phe	Lys	Gly	Trp	Met 1435	Asp	Ile	Met	Tyr	Ala 1440	Ala	Val	Asp
Ser	Arg 1445	Asn	Val	Glu	Leu	Gln 1450	Pro	Lys	Tyr	Glu	Lys 1455	Ser	Leu	Tyr
Met	Tyr 1460		Tyr	Phe	Val	Ile 1465		Ile	Ile	Phe	Gly 1470	Ser	Phe	Phe
Thr	Leu 1475	Asn	Leu	Phe	Ile	Gly 1480	Val	Ile	Ile	Asp	Asn 1485	Phe	Asn	Gln
Gln	Lys 1490	Lys	Lys	Phe	Gly	Gly 1495	Gln	Asp	Ile	Phe	Met 1500	Thr	Glu	Glu
Gln	Lys 1505	Lys	Tyr	Tyr	Asn	Ala 1510	Met	Lys	Lys	Leu	Gly 1515	Ser	Lys	Lys
Pro	Gln 1520	Lys	Pro	Ile	Pro	Arg 1525	Pro	Gly	Asn	Lys	Phe 1530	Gln	Gly	Met

						SCN1	APCT1	ST2	25.tx	ct				
Val	Phe 1535	Asp	Phe	Val	Thr	Arg 1540					Ile 1545		Ile	Met
Ile	Leu 1550	Ile	Cys	Leu	Asn	Met 1555	Val	Thr	Met	Met	Val 1560	Glu	Thr	Asp
Asp	Gln 1 5 65		Glu	Tyr	Val	Thr 1570		Ile	Leu	Ser	Arg 157 5		Asn	Leu
Val	Phe 1580	Ile	Val	Leu	Phe	Thr 1585		Glu	Cys	Val	Leu 1590		Leu	Ile
Ser	Leu 1595		His	Tyr		Phe 1600		Ile	Gly	Trp	Asn 1605		Phe	Asp
Phe	Val 1610	Val	Val	Ile	Leu	Ser 1615	Ile	Val	Gly	Met	Phe 1620	Leu	Ala	Glu
Leu	Ile 1625	Glu	Lys	Tyr	Phe	Val 1630	Ser	Pro	Thr	Leu	Phe 1635	Arg	Val	Ile
Arg	Leu 1640	Ala	Arg	Ile	Gly	Arg 1645	Ile	Leu	Arg	Leu	Ile 1650	Lys	Gly	Ala
-	1655					Leu 1660					1665			
Ala	Leu 1670		Asn	Ile	Gly	Leu 1675		Leu	Phe	Leu	Val 1680	Met	Phe	Ile
Tyr	Ala 1685	Ile	Phe	Gly	Met	Ser 1690	Asn	Phe	Ala	Tyr	Val 1695	Lys	Arg	Glu
Val	Gly 1700	Ile	Asp	Asp	Met	Phe 1705	Asn	Phe	Glu	Thr	Phe 1710	Gly	Asn	Ser
Met	Ile 1715	Cys	Leu	Phe	Gln	Ile 1720	Thr	Thr	Ser	Ala	Gly 1725	Trp	Asp	Gly
Leu	Leu 1730	Ala	Pro	Ile	Leu	Asn 1735	Ser	Lys	Pro	Pro	Asp 1740	Cys	Asp	Pro

SCN1APCT1.ST25.txt

Asn	Lys 1745		Asn	Pro	Gly	Ser 1750		Val	Lys	Gly	Asp 1755		Gly	Asn
Pro	Ser 1760	Val	Gly	Ile	Phe	Phe 1765	Phe	Val	Ser	Tyr	Ile 1770	Ile	Ile	Ser
Phe	Leu 1775	Val	Val	Val	Asn	Met 1780	Tyr	Ile	Ala	Val	Ile 1785	Leu	Glu	Asn
Phe	Ser 1790		Ala	Thr	Glu	Glu 1795	Ser	Ala	Glu	Pro	Leu 1800	Ser	Glu	Asp
Asp	Phe 1805	Glu	Met	Phe	Tyr	Glu 1810	Val	Trp	Glu	Lys	Phe 1815	Asp	Pro	Asp
Ala	Thr 1820	Gln	Phe	Met	Glu	Phe 1825	Glu	Lys	Leu	Ser	Gln 1830	Phe	Ala	Ala
Ala	Leu 1835	Glu	Pro	Pro	Leu	Asn 1840	Leu	Pro	Gln	Pro	Asn 1845	Lys	Leu	Gln
Leu	Ile 1850	Ala	Met	Asp	Leu	Pro 1855	Met	Val	Ser	Gly	Asp 1860	Arg	Ile	His
Cys	Leu 1865	Asp	Ile	Leu	Phe	Ala 1870	Phe	Thr	Lys	Arg	Val 1875	Leu	Gly	Glu
Ser	Gly 1880	Glu	Met	Asp	Ala	Leu 1885	Arg	Ile	Gln	Met	Glu 1890	Glu	Arg	Phe
Met	Ala 1895	Ser	Asn	Pro	Ser	Lys 1900	Val	Ser	Tyr	Gln	Pro 1905	Ile	Ţhr	Thr
Thr	Leu 1910	Lys	Arg	Lys	Gln	Glu 1915	Glu	Val	Ser	Ala	Val 1920	Ile	Ile	Gln
Arg	Ala 1925	Туг	Arg	Arg	His	Leu 1930	Leu	Lys	Arg	Thr	Val 1935	Lys	Gln	Ala
Ser	Phe 1940	Thr	Tyr	Asn	Lys	Asn 1945	Lys	Ile	Lys	Gly	Gly 1950	Ala	Asn	Leu
							Pag	e 99						

SCN1APCT1.ST25.txt

Leu Ile Lys Glu Asp Met Ile Ile Asp Arg Ile Asn Glu Asn Ser 1955 1960 1965

Ile Thr Glu Lys Thr Asp Leu Thr Met Ser Thr Ala Ala Cys Pro 1970 1975 1980

Pro Ser Tyr Asp Arg Val Thr Lys Pro Ile Val Glu Lys His Glu 1985 1990 1995

Gln Glu Gly Lys Asp Glu Lys Ala Lys Gly Lys 2000 2005

<210> 11<211> 8381<212> DNA<213> Homo sapiens<400> 11 atactgcaga ggtctctggt gcatgtgtgt atgtgtgcgt ttgtgtgtgt ttgtgtgtct

60

gtgtgttctg ccccagtgag actgcagccc ttgtaaatac tttgacacct tttgcaagaa 120

ggaatctgaa caattgcaac tgaaggcaca ttgttatcat ctcgtctttg ggtgatgctg
180

ttcctcactg cagatggata attttccttt taatcaggaa tttcatatgc agaataaatg 240

gtaattaaaa tgtgcaggat gacaagatgg agcaaacagt gcttgtacca ccaggacctg 300

acagetteaa ettetteace agagaatete ttgeggetat tgaaagaege attgeagaag 360

aaaaggcaaa gaatcccaaa ccagacaaaa aagatgacga cgaaaatggc ccaaagccaa 420

atagtgactt ggaagctgga aagaaccttc catttattta tggagacatt cctccagaga 480

tggtgtcaga gccctggag gacctggacc cctactatat caataagaaa acttttatag 540

SCN1APCT1.ST25.txt

tattgaataa attgaaggcc atcttccggt tcagtgccac ctctgccctg tacattttaa 600

ctcccttcaa tcctcttagg aaaatagcta ttaagatttt ggtacattca ttattcagca 660

tgctaattat gtgcactatt ttgacaaact gtgtgtttat gacaatgagt aacceteetg
720

attggacaaa gaatgtagaa tacaccttca caggaatata tacttttgaa tcacttataa 780

aaattattgc aaggggattc tgtttagaag attttacttt ccttcgggat ccatggaact

ggctcgattt cactgtcatt acatttgcgt acgtcacaga gtttgtggac ctgggcaatg
900

tctcggcatt gagaacattc agagttctcc gagcattgaa gacgatttca gtcattccag

gcctgaaaac cattgtggga gccctgatcc agtctgtgaa gaagctctca gatgtaatga 1020

tcctgactgt gttctgtctg agcgtatttg ctctaattgg gctgcagctg ttcatgggca
1080

acctgaggaa taaatgtata caatggcctc ccaccaatgc ttccttggag gaacatagta
1140

tagaaaagaa tataactgtg aattataatg gtacacttat aaatgaaact gtctttgagt 1200

ttgactggaa gtcatatatt caagattcaa gatatcatta tttcctggag ggttttttag
1260

atgcactact atgtggaaat agctctgatg caggccaatg tccagaggga tatatgtgtg 1320

tgaaagctgg tagaaatccc aattatggct acacaagctt tgataccttc agttgggctt 1380

SCN1APCT1.ST25.txt

ttttgtcctt gtttcgacta atgactcagg acttctggga aaatctttat caactgacat 1440

tacgtgctgc tgggaaaacg tacatgatat tttttgtatt ggtcattttc ttgggctcat 1500

tctacctaat aaatttgatc ctggctgtgg tggccatggc ctacgaggaa cagaatcagg 1560

ccaccttgga agaagcagaa cagaaagagg ccgaatttca gcagatgatt gaacagctta
1620

aaaagcaaca ggaggcagct cagcaggcag caacggcaac tgcctcagaa cattccagag 1680

agcccagtgc agcaggcagg ctctcagaca gctcatctga agcctctaag ttgagttcca 1740

gggaagagaa agatgaggat gaattccaaa aatctgaatc tgaggacagc atcaggagga 1860

aaggttttcg cttctccatt gaagggaacc gattgacata tgaaaagagg tactcctccc 1920

cacaccagtc tttgttgagc atccgtggct ccctattttc accaaggcga aatagcagaa 1980

caagcetttt cagetttaga gggegageaa aggatgtggg atetgagaae gaettegeag 2040

atgatgagca cagcaccttt gaggataacg agagccgtag agattccttg tttgtgcccc 2100

gacgacacgg agagagacgc aacagcaacc tgagtcagac cagtaggtca tcccggatgc 2160

tggcagtgtt tccagcgaat gggaagatgc acagcactgt ggattgcaat ggtgtggttt

SCN1APCT1.ST25.txt

2220

ccttggttgg tggaccttca gttcctacat cgcctgttgg acagcttctg ccagaggtga 2280

taatagataa gccagctact gatgacaatg gaacaaccac tgaaactgaa atgagaaaga 2340

gaaggtcaag ttctttccac gtttccatgg actttctaga agatccttcc caaaggcaac 2400

gagcaatgag tatagccagc attctaacaa atacagtaga agaacttgaa gaatccaggc 2460

agaaatgccc accetgttgg tataaatttt ccaacatatt cttaatctgg gactgttctc 2520

catatiggtt aaaagtgaaa catgttgtca acctggttgt gatggaccca tttgttgacc 2580

tggccatcac catctgtatt gtcttaaata ctcttttcat ggccatggag cactatccaa 2640

tgacggacca tttcaataat gtgcttacag taggaaactt ggttttcact gggatcttta 2700

cagcagaaat gtttctgaaa attattgcca tggatcctta ctattatttc caagaaggct 2760

ggaatatett tgaeggtttt attgtgaege ttageetggt agaaettgga etegeeaatg 2820

tggaaggatt atctgttctc cgttcatttc gattgctgcg agttttcaag ttggcaaaat 2880

cttggccaac gttaaatatg ctaataaaga tcatcggcaa ttccgtgggg gctctgggaa 2940

atttaaccct cgtcttggcc atcatcgtct tcatttttgc cgtggtcggc atgcagctct 3000

ttggtaaaag ctacaaagat tgtgtctgca agatcgccag tgattgtcaa ctcccacgct
Page 103

SCN1APCT1.ST25.txt

3060

ggcacatgaa tgacttcttc cactccttcc tgattgtgtt ccgcgtgctg tgtggggagt 3120

ggatagagac catgtgggac tgtatggagg ttgctggtca agccatgtgc cttactgtct 3180

tcatgatggt catggtgatt ggaaacctag tggtcctgaa tctctttctg gccttgcttc 3240

tgagctcatt tagtgcagac aaccttgcag ccactgatga tgataatgaa atgaataatc 3300

tccaaattgc tgtggatagg atgcacaaag gagtagctta tgtgaaaaga aaaatatatg 3360

aatttattca acagtccttc attaggaaac aaaagatttt agatgaaatt aaaccacttg 3420

atgatctaaa caacaagaaa gacagttgta tgtccaatca tacaacagaa attgggaaag 3480

atcttgacta tcttaaagat gtaaatggaa ctacaagtgg tataggaact ggcagcagtg 3540

ttgaaaaata cattattgat gaaagtgatt acatgtcatt cataaacaac cccagtctta 3600

ctgtgactgt accaattgct gtaggagaat ctgactttga aaatttaaac acggaagact 3660

ttagtagtga atcggatctg gaagaaagca aagagaaact gaatgaaagc agtagctcat 3720

cagaaggtag cactgtggac atcggcgcac ctgtagaaga acagcccgta gtggaacctg 3780

aagaaactct tgaaccagaa gcttgtttca ctgaaggctg tgtacaaaga ttcaagtgtt 3840

SCN1APCT1.ST25.txt

gtcaaatcaa tgtggaagaa ggcagaggaa aacaatggtg gaacctgaga aggacgtgtt 3900

tccgaatagt tgaacataac tggtttgaga ccttcattgt tttcatgatt ctccttagta
3960

gtggtgctct ggcatttgaa gatatatata ttgatcagcg aaagacgatt aagacgatgt 4020

tggaatatgc tgacaaggtt ttcacttaca ttttcattct ggaaatgctt ctaaaatggg 4080

tggcatatgg ctatcaaaca tatttcacca atgcctggtg ttggctggac ttcttaattg 4140

ttgatgtttc attggtcagt ttaacagcaa atgccttggg ttactcagaa cttggagcca 4200

tcaaatctct caggacacta agagctctga gacctctaag agccttatct cgatttgaag
4260

ggatgagggt ggttgtgaat gcccttttag gagcaattcc atccatcatg aatgtgcttc 4320

tggtttgtct tatattctgg ctaattttca gcatcatggg cgtaaatttg tttgctggca 4380

aattctacca ctgtattaac accacaactg gtgacaggtt tgacatcgaa gacgtgaata 4440

atcatactga ttgcctaaaa ctaatagaaa gaaatgagac tgctcgatgg aaaaatgtga

aagtaaactt tgataatgta ggatttgggt atctctcttt gcttcaagtt gccacattca 4560

aaggatggat ggatataatg tatgcagcag ttgattccag aaatgtggaa ctccagccta 4620

agtatgaaaa aagtetgtae atgtatettt aetttgttat ttteateate tttgggteet 4680

SCN1APCT1.ST25.txt

tetteacett gaacetgttt attggtgtea teatagataa ttteaaceag cagaaaaaga 4740

agtttggagg tcaagacatc tttatgacag aagaacagaa gaaatactat aatgcaatga 4800

aaaaattagg atcgaaaaaa ccgcaaaagc ctatacctcg accaggaaac aaatttcaag 4860

gaatggtett tgaettegta accagacaag tttttgaeat aageateatg atteteatet 4920

gtcttaacat ggtcacaatg atggtggaaa cagatgacca gagtgaatat gtgactacca 4980

ttttgtcacg catcaatctg gtgttcattg tgctatttac tggagagtgt gtactgaaac 5040

tcatctctct acgccattat tattttacca ttggatggaa tatttttgat tttgtggttg 5100

tcattctctc cattgtaggt atgtttcttg ccgagctgat agaaaagtat ttcgtgtccc 5160

ctaccetgtt ecgagtgate egtettgeta ggattggeeg aateetaegt etgateaaag 5220

gagcaaaggg gatccgcacg ctgctctttg ctttgatgat gtcccttcct gcgttgttta
5280

acateggeet ectaetette etagteatgt teatetaege eatetttggg atgteeaact 5340

ttgcctatgt taagagggaa gttgggatcg atgacatgtt caactttgag acctttggca 5400

acagcatgat ctgcctattc caaattacaa cctctgctgg ctgggatgga ttgctagcac 5460

ccattctcaa cagtaagcca cccgactgtg accctaataa agttaaccct ggaagctcag

SCN1APCT1.ST25.txt

5520

ttaagggaga ctgtgggaac ccatctgttg gaattttctt ttttgtcagt tacatcatca

5580

tatectteet ggttgtggtg aacatgtaca tegeggteat eetggagaac tteagtgttg

5640

ctactgaaqa aagtgcagag cctctgagtg aggatgactt tgagatgttc tatgaggttt

5700

gggagaagtt tgatcccgat gcaactcagt tcatggaatt tgaaaaatta tctcagtttg

5760

cagctgcgct tgaaccgcct ctcaatctgc cacaaccaaa caaactccag ctcattgcca

5820

tggatttgcc catggtgagt ggtgaccgga tccactgtct tgatatctta tttgctttta

5880

caaagcgggt tctaggagag agtggagaga tggatgctct acgaatacag atggaagagc

5940

gattcatggc ttccaatcct tccaaggtct cctatcagcc aatcactact actttaaaac

6000

gaaaacaaga ggaagtatet getgteatta tteagegtge ttacagagge cacettttaa

6060

agcqaactqt aaaacaaqct tcctttacqt acaataaaaa caaaatcaaa ggtggggcta

6120

atcttcttat aaaagaagac atgataattg acagaataaa tgaaaactct attacagaaa

6180

aaactgatct gaccatgtcc actgcagctt gtccaccttc ctatgaccgg gtgacaaagc

6240

caattqtqqa aaaacatgag caaqaaggca aaqatqaaaa agccaaaggg aaataaatga

6300

aaataaataa aaataattgg gtgacaaatt gtttacaqcc tgtgaaggtg atgtattttt

SCN1APCT1.ST25.txt

6360

atcaacagga ctcctttagg aggtcaatgc caaactgact gtttttacac aaatctcctt 6420

aaggtcagtg cctacaataa gacagtgacc ccttgtcagc aaactgtgac tctgtgtaaa 6480

ggggagatga ccttgacagg aggttactgt tctcactacc agctgacact gctgaagata 6540

agatgcacaa tggctagtca gactgtaggg accagtttca aggggtgcaa acctgtgatt
6600

ttggggttgt ttaacatgaa acactttagt gtagtaattg tatccactgt ttgcatttca 6660

actgccacat ttgtcacatt tttatggaat ctgttagtgg attcatcttt ttgttaatcc 6720

atgtgtttat tatatgtgac tatttttgta aacgaagttt ctgttgagaa ataggctaag 6780

gacctctata acaggtatgc cacctggggg gtatggcaac cacatggccc tcccagctac . 6840

acaaagtcgt ggtttgcatg agggcatgct gcacttagag atcatgcatg agaaaaagtc 6900

acaagaaaaa caaattetta aattteacea tatttetggg aggggtaatt gggtgataag 6960

tggaggtgct ttgttgatct tgttttgcga aatccagccc ctagaccaag tagattattt
7020

gtgggtaggc cagtaaatct tagcaggtgc aaacttcatt caaatgtttg gagtcataaa 7080

tgttatgttt ctttttgttg tattaaaaaa aaaacctgaa tagtgaatat tgcccctcac 7140

SCN1APCT1.ST25.txt

cctccaccgc cagaagactg aattgaccaa aattactctt tataaatttc tgctttttcc 7200

tgcactttgt ttagccatct ttgggctctc agcaaggttg acactgtata tgttaatgaa 7260

atgctattta ttatgtaaat agtcatttta ccctgtggtg cacgtttgag caaacaaata 7320

atgacctaag cacagtattt attgcatcaa atatgtacca caagaaatgt agagtgcaag 7380

ctttacacag gtaataaaat gtattctgta ccatttatag atagtttgga tgctatcaat 7440

gcatgtttat attaccatgc tgctgtatct ggtttctctc actgctcaga atctcattta 7500

tgagaaacca tatgtcagtg gtaaagtcaa ggaaattgtt caacagatct catttattta
7560

agtcattaag caatagtttg cagcacttta acagcttttt ggttattttt acattttaag 7620

tggataacat atggtatata gccagactgt acagacatgt ttaaaaaaac acactgctta 7680

acctattaaa tatgtgttta gaattttata agcaaatata aatactgtaa aaagtcactt 7740

tattttattt ttcagcatta tgtacataaa tatgaagagg aaattatctt caggttgata 7800

tcacaatcac ttttcttact ttctgtccat agtacttttt catgaaagaa atttgctaaa 7860

taagacatga aaacaagact gggtagttgt agatttctgc tttttaaatt acatttgcta
7920

attttagatt atttcacaat tttaaggagc aaaataggtt cacgattcat atccaaatta 7980

SCN1APCT1.ST25.txt

tgctttgcaa ttggaaaagg gtttaaaatt ttatttatat ttctggtagt acctgtacta 8040

actgaattga aggtagtgct tatgttattt ttgttctttt tttctgactt cggtttatgt 8100

tttcatttct ttggagtaat gctgctctag attgttctaa atagaatgtg ggcttcataa 8160

ttttttttc cacaaaaaca gagtagtcaa cttatatagt caattacatc aggacatttt 8220

gtgtttctta cagaagcaaa ccataggctc ctcttttcct taaaactact tagataaact 8280

gtattcgtga actgcatgct ggaaaatgct actattatgc taaataatgc taaccaacat **

ttaaaatgtg caaaactaat aaagattaca ttttttattt t 8381

Phe Thr Arg Glu Ser Leu Ala Ala Ile Glu Arg Arg Ile Ala Glu Glu 20 25 30

Lys Ala Lys Asn Pro Lys Pro Asp Lys Lys Asp Asp Asp Glu Asn Gly 35 40 45

Pro Lys Pro Asn Ser Asp Leu Glu Ala Gly Lys Asn Leu Pro Phe Ile 50 55 60

Tyr Gly Asp Ile Pro Pro Glu Met Val Ser Glu Pro Leu Glu Asp Leu 65 70 75 80

Asp Pro Tyr Tyr Ile Asn Lys Lys Thr Phe Ile Val Leu Asn Lys Leu 85 90 95

SCN1APCT1.ST25.txt

Lys Ala Ile Phe Arg Phe Ser Ala Thr Ser Ala Leu Tyr Ile Leu Thr Pro Phe Asn Pro Leu Arg Lys Ile Ala Ile Lys Ile Leu Val His Ser Leu Phe Ser Met Leu Ile Met Cys Thr Ile Leu Thr Asn Cys Val Phe Met Thr Met Ser Asn Pro Pro Asp Trp Thr Lys Asn Val Glu Tyr Thr Phe Thr Gly Ile Tyr Thr Phe Glu Ser Leu Ile Lys Ile Ile Ala Arg Gly Phe Cys Leu Glu Asp Phe Thr Phe Leu Arg Asp Pro Trp Asn Trp Leu Asp Phe Thr Val Ile Thr Phe Ala Tyr Val Thr Glu Phe Val Asp Leu Gly Asn Val Ser Ala Leu Arg Thr Phe Arg Val Leu Arg Ala Leu Lys Thr Ile Ser Val Ile Pro Gly Leu Lys Thr Ile Val Gly Ala Leu Ile Gln Ser Val Lys Lys Leu Ser Asp Val Met Ile Leu Thr Val Phe Cys Leu Ser Val Phe Ala Leu Ile Gly Leu Gln Leu Phe Met Gly Asn Leu Arg Asn Lys Cys Ile Gln Trp Pro Pro Thr Asn Ala Ser Leu Glu Glu His Ser Ile Glu Lys Asn Ile Thr Val Asn Tyr Asn Gly Thr Leu Ile Asn Glu Thr Val Phe Glu Phe Asp Trp Lys Ser Tyr Ile Gln Asp

SCN1APCT1.ST25.txt

Ser	Arg	Tyr	His	Tyr 325	Phe	Leu	Glu	Gly	Phe 330	Leu	Asp	Ala	Leu	Leu 335	Cys
Gly	Asn	Ser	Ser 340	Asp	Ala	Gly	Gln	Cys 345	Pro	Glu	Gly	Tyr	Met 350	Cys	Val
Lys	Ala	Gly 355	Arg	Asn	Pro	Asn	Tyr 360	Gly	Tyr	Thr	Ser	Phe 365	Asp	Thr	Phe
Ser	Trp 370	Ala	Phe	Leu	Ser	Leu 375	Phe	Arg	Leu	Met	Thr 380	Gln	Asp	Phe	Trp
Glu 385	Asn	Leu	Tyr	Gln	Leu 390	Thr	Leu	Arg	Ala	Ala 395	Gly	Lys	Thr	Tyr	Met 400
Ile	Phe	Phe	Val	Leu 405	Val	Ile	Phe	Leu	Gly 410	Ser	Phe	Tyr	Leu	Ile 415	Asn
Leu	Ile	Leu	Ala 420	Val	Val	Ala	Met	Ala 425	Tyr	Glu	Glu	Gln	Asn 430	Gln	Ala
Thr	Leu	Glu 435	Glu	Ala	Glu	Gln	Lys 440	Glu	Ala	Glu	Phe	Gln 445	Gln	Met	Ile
Glu	Gln 450	Leu	Lys	Lys	Gln	Gln 455	Glu	Ala	Ala	Gln	Gln 460	Ala	Ala	Thr	Ala
Thr 465	Ala	Ser	Glu	His	Ser 470	Arg	Glu	Pro	Ser	Ala 475	Ala	Gly	Arg	Leu	Ser 480
Asp	Ser	Ser	Ser	Glu 485	Ala	Ser	Lys	Leu	Ser 490	Ser	Lys	Ser	Ala	Lys 495	Glu
Arg	Arg	Asn	Arg 500	Arg	Lys	Lys	Arg	Lys 505	Gln	Lys	Glu	Gln	Ser 510	Gly	Gly
Glu	Glu	Lys 515	Asp	Glu	Asp	Glu	Phe 520	Gln	Lys	Ser	Glu	Ser 525	Glu	Asp	Ser
Ile	Arg	Arg	Lys	Gly	Phe	Arg		Ser ge 1		Glu	Gly	Asn	Arg	Leu	Thr

WO 02/50096	PCT/AU01/01648
-------------	----------------

540

	SCN1APCT1.ST25.txt
530	535

Туr 545	Glu	Lys	Arg	Tyr	Ser 550	Ser	Pro	His	Gln	Ser 555	Leu	Leu	Ser	Ile	Arg 560
Gly	Ser	Leu	Phe	Ser 565	Pro	Arg	Arg	Asn	Ser 570	Arg	Thr	Ser	Leu	Phe 575	Ser
Phe	Arg	Gly	Arg 580	Ala	Lys	Asp	Val	Gly 585	Ser	Glu	Asn	Asp	Phe 590	Ala	Asp
Asp	Glu	His 595	Ser	Thr	Phe	Glu	Asp 600	Asn	Glu	Ser	Arg	Arg 605	Asp	Ser	Leu
Phe	Val 610	Pro	Arg	Arg	His	Gly 615	Glu	Arg	Arg	Asn	Ser 620	Asn	Leu	Ser	Gln
Thr 625	Ser	Arg	Ser	Ser	Arg 630	Met	Leu	Ala	Val	Phe 635	Pro	Ala	Asn	Gly	Lys 640
Met	His	Ser	Thr	Val 645	Asp	Cys	Asn	Gly	Val 650	Val	Ser	Leu	Val	Gly 655	Gly
Pro	Ser	Val	Pro 660	Thr	Ser	Pro	Val	Gly 665	Gln	Leu	Leu	Pro	Glu 670	Val	Ile
Ile	Asp	Lys 675	Pro	Ala	Thr	Asp	Asp 680	Asn	Gly	Thr	Thr	Thr 685	Glu	Thr	Glu
Met	Arg 690	Lys	Arg	Arg	Ser	Ser 695	Ser	Phe	His	Val	Ser 700	Met	Asp	Phe	Leu
Glu 705	Asp	Pro	Ser	Gln	Arg 710	Gln	Arg	Ala	Met	Ser 715	Ile	Ala	Ser	Ile	Leu 720
Thr	Asn	Thr	Val	Glu 725	Glu	Leu	Glu	Glu	Ser 730	Arg	Gln	Lys	Cys	Pro 735	Pro
Cys	Trp	Tyr	Lys 740	Phe	Ser	Asn	Ile	Phe 745	Leu	Ile	Trp	Asp	Cys 750	Ser	Pro

SCN1APCT1.ST25.txt

Tyr Trp Leu Lys Val Lys His Val Val Asn Leu Val Val Met Asp Pro
755 760 765

Phe Val Asp Leu Ala Ile Thr Ile Cys Ile Val Leu Asn Thr Leu Phe 770 775 780

Met Ala Met Glu His Tyr Pro Met Thr Asp His Phe Asn Asn Val Leu 785 790 795 800

Thr Val Gly Asn Leu Val Phe Thr Gly Ile Phe Thr Ala Glu Met Phe 805 810 815

Leu Lys Ile Ile Ala Met Asp Pro Tyr Tyr Tyr Phe Gln Glu Gly Trp 820 825 830

Asn Ile Phe Asp Gly Phe Ile Val Thr Leu Ser Leu Val Glu Leu Gly 835 840 845

Leu Ala Asn Val Glu Gly Leu Ser Val Leu Arg Ser Phe Arg Leu Leu 850 855 860

Arg Val Phe Lys Leu Ala Lys Ser Trp Pro Thr Leu Asn Met Leu Ile 865 870 875 880

Lys Ile Ile Gly Asn Ser Val Gly Ala Leu Gly Asn Leu Thr Leu Val 885 890 895

Leu Ala Ile Ile Val Phe Ile Phe Ala Val Val Gly Met Gln Leu Phe 900 905 910

Gly Lys Ser Tyr Lys Asp Cys Val Cys Lys Ile Ala Ser Asp Cys Gln 915 920 925

Leu Pro Arg Trp His Met Asn Asp Phe Phe His Ser Phe Leu Ile Val 930 935 940

Phe Arg Val Leu Cys Gly Glu Trp Ile Glu Thr Met Trp Asp Cys Met 945 950 955 960

Glu Val Ala Gly Gln Ala Met Cys Leu Thr Val Phe Met Met Val Met 965 970 975

SCN1APCT1.ST25.txt

Val Ile Gly Asn Leu Val Val Leu Asn Leu Phe Leu Ala Leu Leu Leu 980 985 990

Ser Ser Phe Ser Ala Asp Asn Leu Ala Ala Thr Asp Asp Asp Asn Glu 995 1000 1005

Met Asn Asn Leu Gln Ile Ala Val Asp Arg Met His Lys Gly Val 1010 1015 1020

Ala Tyr Val Lys Arg Lys Ile Tyr Glu Phe Ile Gln Gln Ser Phe 1025 1030 1035

Ile Arg Lys Gln Lys Ile Leu Asp Glu Ile Lys Pro Leu Asp Asp 1040 1045 1050

Leu Asn Asn Lys Lys Asp Ser Cys Met Ser Asn His Thr Thr Glu 1055 1060 1065

Ser Gly Ile Gly Thr Gly Ser Ser Val Glu Lys Tyr Ile Ile Asp 1085 1090 1095

Glu Ser Asp Tyr Met Ser Phe Ile Asn Asn Pro Ser Leu Thr Val 1100 1105 1110

Thr Val Pro Ile Ala Val Gly Glu Ser Asp Phe Glu Asn Leu Asn 1115 1120 1125

Thr Glu Asp Phe Ser Ser Glu Ser Asp Leu Glu Glu Ser Lys Glu 1130 1140

Lys Leu Asn Glu Ser Ser Ser Ser Ser Glu Gly Ser Thr Val Asp 1145 1150 1155

Ile Gly Ala Pro Val Glu Glu Gln Pro Val Val Glu Pro Glu Glu 1160 1165 1170

Thr Leu Glu Pro Glu Ala Cys Phe Thr Glu Gly Cys Val Gln Arg 1175 1180 1185

SCN1APCT1.ST25.txt

Phe	Lys 1190		Cys	Gln	Ile	Asn 1195		Glu	Glu	Gly	Arg 1200	Gly	Lys	Gln
Trp	Trp 1205	Asn	Leu	Arg	Arg	Thr 1210		Phe	Arg	Ile	Val 1215	Glu	His	Asn
Trp	Phe 1220	Glu	Thr	Phe	Ile	Val 1225		Met	Ile	Leu	Leu 1230	Ser	Ser	Gly
Ala	Leu 1235	Ala	Phe	Glu	Asp	Ile 1240	Tyr	Ile	Asp	Gln	Arg 1245	Lys	Thr	Ile
Lys	Thr 1250	Met	Leu	Glu	Tyr	Ala 1255	Asp	Lys	Val	Phe	Thr 1260	Tyr	Ile	Phe
Ile	Leu 1265	Glu	Met	Leu	Leu	Lys 1270		Val	Ala	Tyr	Gly 1275	Tyr	Gln	Thr
Tyr	Phe 1280	Thr	Asn	Ala	Trp	Cys 1285		Leu	Asp	Phe	Leu 1290	Ile	Val	Asp
Val	Ser 1295	Leu	Val	Ser	Leu	Thr 1300	Ala	Asn	Ala	Leu	Gly 1305	Tyr	Ser	Glu
Leu	Gly 1310	Ala	Ile	Lys	Ser	Leu 1315	Arg	Thr	Leu	Arg	Ala 1320	Leu	Arg	Pro
Leu	Arg 1325	Ala	Leu	Ser	Arg	Phe 1330	G1u	Gly	Met	Arg	Va1 1335	Val	Val	Asn
Ala	Leu 1340	Leu	Gly	Ala	Ile	Pro 1345	Ser	Ile	Met	Asn	Val 1350	Leu	Leu	Val
Cys	Leu 1355	Ile	Phe	Trp	Leu	I1e 1360	Phe	Ser	Ile	Met	Gly 1365	Val	Asn	Leu
Phe	Ala 1370	Gly	Lys	Phe	Tyr	His 1375	Cys	Ile	Asn	Thr	Thr 1380	Thr	Gly	Asp
Arg	Phe	Asp	Ile	Glu	Asp	Val		Asn 116		Thr	Asp	Cys	Leu	Lys

1395

SCN1APCT1.ST25.txt 1385 1390

Leu	Ile 1400	Glu	Arg	Asn	Glu	Thr 1405	Ala	Arg	Trp	Lys	Asn 1410	Val	Lys	Val
Asn	Phe 1415	Asp	Asn	Val	Gly	Phe 1420	Gly	Tyr	Leu	Ser	Leu 1425	Leu	Gln	Val
Ala	Thr 1430		Lys	Gly	Trp	Met 1435	Asp	Ile	Met	Tyr	Ala 1440	Ala	Val	Asp
Ser	Arg 1445		Val	Glu	Leu	Gln 1450	Pro	Lys	Tyr	Glu	Lys 1455	Ser	Leu	Tyr
Met	Tyr 1460		Tyr	Phe	Val	Ile 1465		Ile	Ile	Phe	Gly 1470	Ser	Phe	Phe
Thr	Leu 1475	Asn	Leu	Phe	Ile	Gly 1480	Val	Ile	Ile	Asp	Asn 1485	Phe	Asn	Gln
Gln	Lys 1490		Lys	Phe	Gly	Gly 1495	Gln	Asp	Ile	Phe	Met 1500	Thr	Glu	Glu
Gln	Lys 1505	Lys	Tyr	Tyr	Asn	Ala 1510	Met	Lys	Lys	Leu	Gly 1515	Ser	Lys	Lys
Pro	Gln 1520		Pro	Ile	Pro	Arg 1525	Pro	Gly	Asn	Lys	Phe 1530	Gln	Gly	Met
	Phe 1535	Asp	Phe	Val		Arg 1540		Val	Phe		Ile 1545		Ile	Met
Ile	Leu 1550	Ile	Cys	Leu	Asn	Met 1555	Val	Thr	Met	Met	Val 1560	Glu	Thr	Asp
Asp	Gln 1565	Ser	Glu	Tyr	Val	Thr 1570	Thr	Ile	Leu	Ser	Arg 1575	Ile	Asn	Leu
Val	Phe 1580	Ile	Val	Leu	Phe	Thr 1585	Gly	Glu	Cys	Val	Leu 1590	Lys	Leu	Ile

Ser	Leu 1595	Arg	His	Tyr		SCN1A Phe 1600	Thr				Asn 1605	Ile	Phe	Asp
Phe	Val 1610		Val	Ile	Leu	Ser 1615	Ile	Val	Gly	Met	Phe 1620		Ala	Glu
Leu	Ile 1625	Glu	Lys	Tyr	Phe	Val 1630	Ser	Pro	Thr	Leu	Phe 1635	Arg	Val	Ile
Arg	Leu 1640	Ala	Arg	Ile	Gly	Arg 1645	Ile	Leu	Arg	Leu	Ile 1650	Lys	Gly	Ala
Lys	Gly 1655	Ile	Arg	Thr	Leu	Leu 1660	Phe	Ala	Leu	Met	Met 1665	Ser	Leu	Pro
Ala	Leu 1670	Phe	Asn	Ile	Gly	Leu 1675	Leu	Leu	Phe	Leu	Val 1680	Met	Phe	Ile
Tyr	Ala 1685		Phe	Gly	Met	Ser 1690	Asn	Phe	Ala	Tyr	Val 1695	Lys	Arg	Glu
Val	Gly 1700	Ile	Asp	Asp	Met	Phe 1705	Asn	Phe	Glu	Thr	Phe 1710	Gly	Asn	Ser
Met	Ile 1715		Leu	Phe	Gln	Ile 1720	Thr	Thr	Ser	Ala	Gly 1725	Trp	Asp	Gly
Leu	Leu 1730	Ala	Pro	Ile	Leu	Asn 1735	Ser	Lys	Pro	Pro	Asp 1740	Cys	Asp	Pro
Asn	Lys 1745	Val	Asn	Pro	Gly	Ser 1 7 50	Ser	Val	Lys	Gly	Asp 1755	Суѕ	Gly	Asn
Pro	Ser 1760	Val	Gly	Ile	Phe	Phe 1765	Phe	Val	Ser	Tyr	Ile 1770	Ile	Ile	Ser
Phe	Leu 17 7 5	Val	Val	Val	Asn	Met 1780	Tyr	Ile	Ala	Val	Ile 1785	Leu	Glu	Asn
Phe	Ser 1 7 90	Val	Ala	Thr	Glu	Glu 1795	Ser	Ala	Glu	Pro	Leu 1800	Ser	Gl u	Asp

SCN1APCT1.ST25.txt

Asp	Phe 1805	Glu	Met	Phe	Tyr	Glu 1810	Val	Trp	Glu	Lys	Phe 1815	Asp	Pro	Asp
Ala	Thr 1820	Gln	Phe	Met	Glu	Phe 1825	Glu	Lys	Leu	Ser	Gln 1830	Phe	Ala	Ala
Ala	Leu 1835		Pro	Pro	Leu	Asn 1840		Pro	Gln	Pro	Asn 1845	_	Leu	Gln
Leu	Ile 1850	Ala	Met	Asp	Leu	Pro 1855	Met	Val	Ser	Gly	Asp 1860	Arg	Ile	His
Cys	Leu 1865	Asp	Ile	Leu	Phe	Ala 1870	Phe	Thr	Lys	Arg	Val 1875	Leu	Gly	Glu
Ser	Gly 1880	Glu	Met	Asp	Ala	Leu 1885	Arg	Ile	Gln	Met	Glu 1890	Glu	Arg	Phe
Met	Ala 1895	Ser	Asn	Pro	Ser	Lys 1900	Val	Ser	Tyr	Gln	Pro 1905	Ile	Thr	Thr
Thr	Leu 1910	Lys	Arg	Lys	Gln	Glu 1915	Glu	Val	Ser	Ala	Val 1920	Ile	Ile	Gln
Arg	Ala 1925	Tyr	Arg	Gly	His	Leu 1930	Leu	Lys	Arg	Thr	Val 1935	Lys	Gln	Ala
Ser	Phe 1940	Thr	Tyr	Asn	Lys	Asn 1945	Lys	Ile	Lys	Gly	Gly 1950	Ala	Asn	Leu
Leu	Ile 1955	Lys	Glu	Asp	Met	Ile 1960	Ile	Asp	Arg	Ile	Asn 1965	Glu	Asn	Ser
Ile	Thr 1970	Glu	Lys	Thr	Asp	Leu 1975	Thr	Met	Ser	Thr	Ala 1980	Ala	Cys	Pro
Pro	Ser 1985	Tyr	Asp	Arg	Val	Thr 1990	Lys	Pro	Ile	Val	Glu 1995	Lys	His	Glu
Gln	Glu 2000	Gly	Lys	Asp	Glu	Lys 2005	Ala	_	_	Lys				

SCN1APCT1.ST25.txt

<210> 13<211> 27<212> DNA<213> agatgaccag agtgaatatg tgactac	Homo sapiens<400>	13
<210> 14<211> 24<212> DNA<213> ccaatggtaa aataataatg gcgt	Homo sapiens<400>	14
24		
<210> 15<211> 20<212> DNA<213> taccatagag tgaggcgagg	Homo sapiens<400>	15
20		
<210> 16<211> 20<212> DNA<213> atggacttcc tgctctgccc	Homo sapiens<400>	16
20		
<210> 17<211> 22<212> DNA<213> cctctagctc atgtttcatg ac	Homo sapiens<400>	17
22		
<210> 18<211> 20<212> DNA<213> tgcagtaggc aattagcagc	Homo sapiens<400>	18
20		
<210> 19<211> 26<212> DNA<213> ctaattaaga agagatccag tgacag	Homo sapiens<400>	19
26		
<210> 20<211> 27<212> DNA<213> gctataaagt gcttacagat catgtac	Homo sapiens<400>	20
27		
<210> 21<211> 24<212> DNA<213> ccctgaattt tggctaagct gcag	Homo sapiens<400>	21
24		

SCN1APCT1.ST25.txt

<210> 22<211> 27<212> ctacattaag acacagtttc aa		Homo	sapiens<400>	22
27				
<210> 23<211> 21<212> gggctacgtt tcatttgtat g	DNA<213>	Homo	sapiens<400>	23
21				
<210> 24<211> 27<212> gcaacctatt cttaaagcat aag		Homo	sapiens<400>	24
27				
<210> 25<211> 20<212> aggctctttg tacctacagc	DNA<213>	Homo	sapiens<400>	25
20				
<210> 26<211> 20<212> catgtagggt ccgtctcatt	DNA<213>	Homo	sapiens<400>	26
20				
<210> 27<211> 23<212> cacacgtgtt aagtcttcat agt		Homo	sapiens<400>	27
23				
<210> 28<211> 21<212> agcccctcaa gtatttatcc t	DNA<213>	Homo	sapiens<400>	28
21				
<210> 29<211> 21<212> gaacctgacc ttcctgttct c	DNA<213>	Homo	sapiens<400>	29
21				
<210> 30<211> 22<212> gttggctgtt atcttcagtt tc	DNA<213>	Homo	sapiens<400>	30
22				

<210> 31<211> 23<212> gactaggcaa tatcatagca ta			.txt sapiens<400>	31
23				
<210> 32<211> 23<212> ctttctacta tattatcatc cg		Homo	sapiens<400>	32
23				
<210> 33<211> 20<212> ttgaaagttg aagccaccac	DNA<213>	Homo	sapiens<400>	33
<210> 34<211> 20<212> ccacctgctc ttaggtactc	DNA<213>	Homo	sapiens<400>	34
20				
<210> 35<211> 21<212> gccatgcaaa tacttcagcc c	DNA<213>	Homo	sapiens<400>	35
<210> 36<211> 23<212> cacaacagtg gttgattcag tt		Homo	sapiens<400>	36
<pre>23 <210> 37<211> 23<212> tgaatgctga aatctccttc ta</pre>		Homo	sapiens<400>	37
23				
<210> 38<211> 21<212> ctcaggttgc tgttgcgtct c	DNA<213>	Homo	sapiens<400>	38
21				
<210> 39<211> 21<212> gataacgaga gccgtagaga t	DNA<213>	Homo	sapiens<400>	39
21				
<210> 40<211> 20<212>	DNA<213> Page		sapiens<400>	40

tctgtagaaa cactggctgg

20

<210> 41<211> 21<212> DNA<213> Homo sapiens<400> 41 catgaaattc actgtgtcac c

21

<210> 42<211> 21<212> DNA<213> Homo sapiens<400> 42 cagctcttga attagactgt c

21

<210> 43<211> 20<212> DNA<213> Homo sapiens<400> 43 atccttggga ggtttagagt

20

<210> 44<211> 21<212> DNA<213> Homo sapiens<400> 44 catcacaacc aggttgacaa c

21

<210> 45<211> 21<212> DNA<213> Homo sapiens<400> 45 ctgggactgt tctccatatt g

21

<210> 46<211> 20<212> DNA<213> Homo sapiens<400> 46 gcatgaagga tggttgaaag

20

<210> 47<211> 23<212> DNA<213> Homo sapiens<400> 47 cattgtggga aaatagcata agc

23

<210> 48<211> 20<212> DNA<213> Homo sapiens<400> 48 gctatgcaga accctgattg

20

<210> 49<211> 20<212> DNA<213> Homo sapiens<400> 49 tgagacggtt agggcagatc

SCN1APCT1.ST25.txt

20

<210> 50<211> 21<212> agaagtcatt catgtgccag c	DNA<213>	Homo	sapiens<400>	50
21 <210> 51<211> 21<212> ctgcaagatc gccagtgatt g	DNA<213>	Homo	sapiens<400>	51
21				
<210> 52<211> 20<212> acatgtgcac aatgtgcagg 20	DNA<213>	Homo	sapiens<400>	52
<210> 53<211> 22<212> gtggtgtttc cttctcatca ag	DNA<213>	Homo	sapiens<400>	53
<pre>22 <210> 54<211> 22<212> tctgctgtat gattggacat ac</pre>	DNA<213>	Homo	sapiens<400>	54
22	DNIA - 0.1.2.	Home	ani ong (100)	55
<pre><210> 55<211> 22<212> caacagtcct tcattaggaa ac 22</pre>	DNA<513>	ношо	sapiens<400>	22
<210> 56<211> 22<212> accttcccac acctatagaa tc	DNA<213>	Homo	sapiens<400>	56
<210> 57<211> 21<212> cttggcaggc aacttattac c	DNA<213>	Homo	sapiens<400>	57
21				5 0
<pre><210> 58<211> 22<212> caagctgcac tccaaatgaa ag</pre>	DNA<213>	Homo	sapiens<400>	58

22

 $<\!210\!>$ 59<211> 24<212> DNA<213> Homo sapiens<400> 59 tggaagcaga gacactttat ctac

24

<210> 60<211> 24<212> DNA<213> Homo sapiens<400> 60 gtgctgtatc accttttctt aatc

24

<210> 61<211> 24<212> DNA<213> Homo sapiens<400> 61 cctattccaa tgaaatgtca tatg

24

<210> 62<211> 21<212> DNA<213> Homo sapiens<400> 62 caagetacet tgaacagaga c

21

<210> 63<211> 24<212> DNA<213> Homo sapiens<400> 63 ctacacattg aatgatgatt ctgt

24

<210> 64<211> 24<212> DNA<213> Homo sapiens<400> 64 gctatataca atacttcagg ttct

24

<210> 65<211> 21<212> DNA<213> Homo sapiens<400> 65 accagagatt actaggggaa t

21

<210> 66<211> 21<212> DNA<213> Homo sapiens<400> 66 ccatcgagca gtctcatttc t

21

<210> 67<211> 21<212> DNA<213> Homo sapiens<400> 67 acaactggtg acaggtttga c

21

SCN1APCT1.ST25.txt

<210> 68<211> 24<212> DNA <ctgggctcat aaacttgtac="" taac<="" th=""><th>213> Homo sapiens<400></th><th>> 68</th></ctgggctcat>	213> Homo sapiens<400>	> 68
24		
<210> 69<211> 21<212> DNA< actgtcttgg tccaaaatct g	213> Homo sapiens<400>	> 69
21		
<210> 70<211> 24<212> DNA< ttcgattaat tttaccacct gatc	213> Homo sapiens<400>	· 70
24		
<210> 71<211> 21<212> DNA< agcaccagtg acatttccaa c	213> Homo sapiens<400>	> 71
21		
<210> 72<211> 21<212> DNA< ggcagagaaa acactccaag g	213> Homo sapiens<400>	> 72
21		
<210> 73<211> 21<212> DNA< gacacagttt taaccagttt g	213> Homo sapiens<400>	> 73
21		
<210> 74<211> 21<212> DNA< tgtgagacaa gcatgcaagt t	213> Homo sapiens<400>	> 74
21		
<210> 75<211> 21<212> DNA< cagggccaat gactactttg c	213> Homo sapiens<400>	> 75
21		
<210> 76<211> 25<212> DNA< ctgattgctg ggatgatctt gaatc	213> Homo sapiens<400>	→ 76
25		

SCN1APCT1.ST25.txt

SCNIAPCTI.ST25.CXC				
<210> 77<211> 23<212> cgcatgattt cttcactggt tgg	•	Homo	sapiens<400>	77
23				
<210> 78<211> 21<212> gcgtagatga acatgactag g	DNA<213>	Homo	sapiens<400>	78
21				
<210> 79<211> 21<212> tcctgcgttg tttaacatcg g	DNA<213>	Homo	sapiens<400>	79
21				
<210> 80<211> 21<212> attccaacag atgggttccc a	DNA<213>	Homo	sapiens<400>	80
21				
<210> 81<211> 21<212> tggaagctca gttaagggag a	DNA<213>	Homo	sapiens<400>	81
21				
<210> 82<211> 21<212> agcgcagctg caaactgaga t	DNA<213>	Homo	sapiens<400>	82
21				
<210> 83<211> 22<212> ccgatgcaac tcagttcatg ga	DNA<213>	Homo	sapiens<400>	83
22				
<210> 84<211> 21<212> gtagtgattg gctgatagga g	DNA<213>	Homo	sapiens<400>	84
21				
<210> 85<211> 24<212> agagcgattc atggcttcca atc		Homo	sapiens<400>	85

Page 127

24

SCN1APCT1.ST25.txt

<210> 86<211> 26<212> DNA<213> Homo sapiens<400> 86 tgccttcttg ctcatgtttt tccaca

26

<210> 87<211> 22<212> DNA<213> Homo sapiens<400> 87 cctatgaccg ggtgacaaag cc

22

<210> 88<211> 22<212> DNA<213> Homo sapiens<400> 88 tgctgacaag gggtcactgt ct

22

<210> 89<211> 8381<212> DNA<213> Homo sapiens<400> 89 atactgcaga ggtctctggt gcatgtgtgt atgtgtgcgt ttgtgtgtgt ttgtgtgtct

60

gtgtgttctg ccccagtgag actgcagccc ttgtaaatac tttgacacct tttgcaagaa 120

ggaatctgaa caattgcaac tgaaggcaca ttgttatcat ctcgtctttg ggtgatgctg
180

ttcctcactg cagatggata attttccttt taatcaggaa tttcatatgc agaataaatg 240

gtaattaaaa tgtgcaggat gacaagatgg agcaaacagt gcttgtacca ccaggacctg

acagetteaa ettetteaee agagaatete ttgeggetat tgaaagaege attgeagaag 360

aaaaggcaaa gaatcccaaa ccagacaaaa aagatgacga cgaaaatggc ccaaagccaa 420

atagtgactt ggaagctgga aagaaccttc catttattta tggagacatt cctccagaga 480

tggtgtcaga gccctggag gacctggacc cctactatat caataagaaa acttttatag 540

SCN1APCT1.ST25.txt

tattgaataa attgaaggcc atcttccggt tcagtgccac ctctgccctg tacattttaa 600

ctcccttcaa tcctcttagg aaaatagcta ttaagatttt ggtacattca ttattcagca 660

tgctaattat gtgcactatt ttgacaaact gtgtgtttat gacaatgagt aaccctcctg
720

attggacaaa gaatgtagaa tacaccttca caggaatata tacttttgaa tcacttataa 780

aaattattgc aaggggattc tgtttagaag attttacttt ccttcgggat ccatggaact 840

ggctcgattt cactgtcatt acatttgcgt acgtcacaga gtttgtggac ctgggcaatg
900

tctcggcatt gagaacattc agagttctcc gagcattgaa gacgatttca gtcattccag
960

gcctgaaaac cattgtggga gccctgatcc agtctgtgaa gaagctctca gatgtaatga
1020

tectgactgt gttetgtetg agegtatttg etetaattgg getgeagetg tteatgggea
1080

acctgaggaa taaatgtata caatggcctc ccaccaatgc ttccttggag gaacatagta 1140

tagaaaagaa tataactgtg aattataatg gtacacttat aaatgaaact gtctttgagt 1200

ttgactggaa gtcatatatt caagattcaa gatatcatta tttcctggag ggttttttag
1260

atgcactact atgtggaaat agctctgatg caggccaatg tccagaggga tatatgtgtg
1320

tgaaagctgg tagaaatccc aattatggct acacaagctt tgataccttc agttgggctt 1380

Page 129

SCN1APCT1.ST25.txt

ttttgtcctt gtttcgacta atgactcagg acttctggga aaatctttat caactgacat 1440

tacgtgctgc tgggaaaacg tacatgatat tttttgtatt ggtcattttc ttgggctcat 1500

tctacctaat aaatttgatc ctggctgtgg tggccatggc ctacgaggaa cagaatcagg
1560

ccaccttgga agaagcagaa cagaaagagg ccgaatttca gcagatgatt gaacagctta

1620

aaaagcaaca ggaggcagct cagcaggcag caacggcaac tgcctcagaa cattccagag

agcccagtgc agcaggcagg ctctcagaca gctcatctga agcctctaag ttgagttcca 1740

gggaagagaa agatgaggat gaattccaaa aatctgaatc tgaggacagc atcaggagga 1860

aaggttttcg cttctccatt gaagggaacc gattgacata tgaaaagagg tactcctccc 1920

cacaccagtc tttgttgagc atccgtggct ccctattttc accaaggcga aatagcagaa 1980

caageetttt cagetttaga gggegageaa aggatgtggg atetgagaac gaettegeag 2040

atgatgagca cagcaccttt gaggataacg agagccgtag agattccttg tttgtgcccc 2100

gacgacacgg agagagacgc aacagcaacc tgagtcagac cagtaggtca tcccggatgc 2160

tggcagtgtt tccagcgaat gggaagatgc acagcactgt ggattgcaat ggtgtggttt

SCN1APCT1.ST25.txt

2220

ccttggttgg tggaccttca gttcctacat cgcctgttgg acagcttctg ccagaggtga 2280

taatagataa gccagctact gatgacaatg gaacaaccac tgaaactgaa atgagaaaga 2340

gaaggtcaag ttctttccac gtttccatgg actttctaga agatccttcc caaaggcaac 2400

gagcaatgag tatagccagc attctaacaa atacagtaga agaacttgaa gaatccaggc 2460

agaaatgccc accetgttgg tataaatttt ccaacatatt cttaatctgg gactgttctc 2520

catattggtt aaaagtgaaa catgttgtca acctggttgt gatggaccca tttgttgacc 2580

tggccatcac catctgtatt gtcttaaata ctcttttcat ggccatggag cactatccaa 2640

tgacggacca tttcaataat gtgcttacag taggaaactt ggttttcact gggatcttta 2700

cagcagaaat gtttctgaaa attattgcca tggatcctta ctattatttc caagaaggct 2760

ggaatatett tgaeggtttt attgtgaege ttageetggt agaaettgga etegeeaatg 2820

tggaaggatt atctgttctc cgttcatttc gattgctgcg agttttcaag ttggcaaaat 2880

cttggccaac gttaaatatg ctaataaaga tcatcggcaa ttccgtgggg gctctgggaa 2940

atttaaccct cgtcttggcc atcatcgtct tcatttttgc cgtggtcggc atgcagctct 3000

ttggtaaaag ctacaaagat tgtgtctgca agatcgccag tgattgtcaa ctcccacgct
Page 131

SCN1APCT1.ST25.txt

3060

ggcacatgaa tgacttcttc cactccttcc tgattgtgtt ccgcgtgctg tgtggggagt 3120

ggatagagac catgtgggac tgtatggagg ttgctggtca agccatgtgc cttactgtct 3180

tcatgatggt catggtgatt ggaaacctag tggtcctgaa tctctttctg gccttgcttc 3240

tgagctcatt tagtgcagac aaccttgcag ccactgatga tgataatgaa atgaataatc . 3300

tccaaattgc tgtggatagg atgcacaaag gagtagctta tgtgaaaaga aaaatatatg 3360

aatttattca acagtccttc attaggaaac aaaagatttt agatgaaatt aaaccacttg 3420

atgatctaaa caacaagaaa gacagttgta tgtccaatca tacaacagaa attgggaaag 3480

atcttgacta tcttaaagat gtaaatggaa ctacaagtgg tataggaact ggcagcagtg 3540

ttgaaaaata cattattgat gaaagtgatt acatgtcatt cataaacaac cccagtctta 3600

ctgtgactgt accaattgct gtaggagaat ctgactttga aaatttaaac acggaagact 3660

ttagtagtga atcggatctg gaagaaagca aagagaaact gaatgaaagc agtagctcat 3720

cagaaggtag cactgtggac atcggcgcac ctgtagaaga acagcccgta gtggaacctg 3780

aagaaactct tgaaccagaa gcttgtttca ctgaaggctg tgtacaaaga ttcaagtgtt 3840

SCN1APCT1.ST25.txt

gtcaaatcaa tgtggaagaa ggcagaggaa aacaatggtg gaacctgaga aggacgtgtt 3900

tccgaatagt tgaacataac tggtttgaga ccttcattgt tttcatgatt ctccttagta 3960

gtggtgctct ggcatttgaa gatatatata ttgatcagcg aaagacgatt aagacgatgt 4020

tggaatatgc tgacaaggtt ttcacttaca ttttcattct ggaaatgctt ctaaaatggg 4080

tggcatatgg ctatcaaaca tatttcacca atgcctggtg ttggctggac ttcttaattg 4140

ttgatgtttc attggtcagt ttaacagcaa atgccttggg ttactcagaa cttggagcca 4200

tcaaatctct caggacacta agagctctga gacctctaag agccttatct cgatttgaag 4260

ggatgagggt ggttgtgaat gcccttttag gagcaattcc atccatcatg aatgtgcttc 4320

tggtttgtct tatattctgg ctaattttca gcatcatggg cgtaaatttg tttgctggca . 4380

aattctacca ctgtattaac accacaactg gtgacaggtt tgacatcgaa gacgtgaata 4440

atcatactga ttgcctaaaa ctaatagaaa gaaatgagac tgctcgatgg aaaaatgtga 4500

aagtaaactt tgataatgta ggatttgggt atctctcttt gcttcaagtt gccacattca 4560

aaggatggat ggatataatg tatgcagcag ttgattccag aaatgtggaa ctccagccta

agtatgaaaa aagtctgtac atgtatcttt actttgttat tttcatcatc tttgggtcct 4680

SCN1APCT1.ST25.txt

tottcacctt gaacctgttt attggtgtca tcatagataa tttcaaccag cagaaaaaga 4740

agtttggagg tcaagacatc tttatgacag aagaacagaa gaaatactat aatgcaatga 4800

aaaaattagg atcgaaaaaa ccgcaaaagc ctatacctcg accaggaaac aaatttcaag
4860

gaatggtett tgaettegta accagacaag tttttgaeat aageateatg atteteatet 4920

gtcttaacat ggtcacaatg atggtggaaa cagatgacca gagtgaatat gtgactacca 4980

ttttgtcacg catcaatctg gtgttcattg tgctatttac tggagagtgt gtactgaaac 5040

tcatctctct acgccattat tattttacca ttggatggaa tatttttgat tttgtggttg 5100

tcattctctc cattgtaggt atgtttcttg ccgagctgat agaaaagtat ttcgtgtccc 5160

ctaccetgtt eegagtgate egtettgeta ggattggeeg aateetaegt etgateaaag 5220

gagcaaaggg gatccgcacg ctgctctttg ctttgatgat gtcccttcct gcgttgttta
5280

acateggeet ectaetette etagteatgt teatetaege eatetttggg atgteeaaet 5340

ttgcctatgt taagagggaa gttgggatcg atgacatgtt caactttgag acctttggca 5400

acagcatgat ctgcctattc caaattacaa cctctgctgg ctgggatgga ttgctagcac 5460

ccattctcaa cagtaagcca cccgactgtg accctaataa agttaaccct ggaagctcag

SCN1APCT1.ST25.txt

5520

ttaagggaga ctgtgggaac ccatctgttg gaattttctt ttttgtcagt tacatcatca 5580

tatectteet ggttgtggtg aacatgtaca tegeggteat eetggagaac tteagtgttg 5640

ctactgaaga aagtgcagag cctctgagtg aggatgactt tgagatgttc tatgaggttt 5700

gggagaagtt tgatcccgat gcaactcagt tcatggaatt tgaaaaatta tctcagtttg 5760

cagctgcgct tgaaccgcct ctcaatctgc cacaaccaaa caaactccag ctcattgcca 5820

tggatttgcc catggtgagt ggtgaccgga tccactgtct tgatatctta tttgctttta
5880

caaagcgggt tctaggagag agtggagaga tggatgctct acgaatacag atggaagagc 5940

gattcatggc ttccaatcct tccaaggtct cctatcagcc aatcactact actttaaaac 6000

gaaaacaaga ggaagtatct gctgtcatta ttcagcgtgc ttacagacgc caccttttaa 6060

agcgaactgt aaaacaagct tcctttacgt acaataaaaa caaaatcaaa ggtggggcta 6120

atcttcttat aaaagaagac atgataattg acagaataaa tgaaaactct attacagaaa 6180

aaactgatct gaccatgtcc actgcagctt gtccaccttc ctatgaccgg gtgacaaagc 6240

caattgtgga aaaacatgag caagaaggca aagatgaaaa agccaaaggg aaataaatga 6300

aaataaataa aaataattgg gtgacaaatt gtttacagcc tgtgaaggtg atgtattttt
Page 135

SCN1APCT1.ST25.txt

6360

atcaacagga ctcctttagg aggtcaatgc caaactgact gtttttacac aaatctcctt 6420

aaggtcagtg cctacaataa gacagtgacc ccttgtcagc aaactgtgac tctgtgtaaa 6480

ggggagatga ccttgacagg aggttactgt tctcactacc agctgacact gctgaagata 6540

agatgcacaa tggctagtca gactgtaggg accagtttca aggggtgcaa acctgtgatt 6600

ttggggttgt ttaacatgaa acactttagt gtagtaattg tatccactgt ttgcatttca 6660

actgccacat ttgtcacatt tttatggaat ctgttagtgg attcatcttt ttgttaatcc 6720

atgtgtttat tatatgtgac tatttttgta aacgaagttt ctgttgagaa ataggctaag 6780

gacctctata acaggtatgc cacctggggg gtatggcaac cacatggccc tcccagctac 6840

acaaagtcgt ggtttgcatg agggcatgct gcacttagag atcatgcatg agaaaaagtc 6900

acaagaaaaa caaattetta aattteacea tatttetggg aggggtaatt gggtgataag 6960

tggaggtgct ttgttgatct tgttttgcga aatccagccc ctagaccaag tagattattt 7020

gtgggtaggc cagtaaatct tagcaggtgc aaacttcatt caaatgtttg gagtcataaa 7080

tgttatgttt ctttttgttg tattaaaaaa aaaacctgaa tagtgaatat tgcccctcac 7140

SCN1APCT1.ST25.txt

cctccaccgc cagaagactg aattgaccaa aattactctt tataaatttc tgctttttcc 7200

tgcactttgt ttagccatct ttgggctctc agcaaggttg acactgtata tgttaatgaa 7260

atgctattta ttatgtaaat agtcatttta ccctgtggtg cacgtttgag caaacaaata 7320

atgacctaag cacagtattt attgcatcaa atatgtacca caagaaatgt agagtgcaag 7380

ctttacacag gtaataaaat gtattctgta ccatttatag atagtttgga tgctatcaat 7440

gcatgtttat attaccatgc tgctgtatct ggtttctctc actgctcaga atctcattta
7500

tgagaaacca tatgtcagtg gtaaagtcaa ggaaattgtt caacagatct catttattta
7560

agtcattaag caatagtttg cagcacttta acagcttttt ggttattttt acattttaag 7620

tggataacat atggtatata gccagactgt acagacatgt ttaaaaaaac acactgctta
7680

acctattaaa tatgtgttta gaattttata agcaaatata aatactgtaa aaagtcactt 7740

tattttattt ttcagcatta tgtacataaa tatgaagagg aaattatctt caggttgata 7800

tcacaatcac ttttcttact ttctgtccat agtacttttt catgaaagaa atttgctaaa 7860

taagacatga aaacaagact gggtagttgt agatttctgc tttttaaatt acatttgcta
7920

attttagatt atttcacaat tttaaggagc aaaataggtt cacgattcat atccaaatta 7980

SCN1APCT1.ST25.txt

tgctttgcaa ttggaaaagg gtttaaaatt ttatttatat ttctggtagt acctgtacta 8040

actgaattga aggtagtgct tatgttattt ttgttctttt tttctgactt cggtttatgt 8100

tttcatttct ttggagtaat gctgctctag attgttctaa atagaatgtg ggcttcataa 8160

ttttttttc cacaaaaaca gagtagtcaa cttatatagt caattacatc aggacatttt 8220

gtgtttctta cagaagcaaa ccataggctc ctcttttcct taaaactact tagataaact 8280

gtattcgtga actgcatgct ggaaaatgct actattatgc taaataatgc taaccaacat 8340

ttaaaatgtg caaaactaat aaagattaca tttttattt t 8381

International application No.

PCT/AU01/01648

A. CLASSIFICATION OF SUBJECT MATTER

Int. Cl. 7: C07H 21/04; C07K 14/435, 16/18; C12N 15/12, 15/63; A61K 38/17, 39/395, 31/7105, 48/00; A61P 25/08

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) STN search in CA, Medline, WPIDS, BIOSIS. Keywords: sodium channel, mutat?, epilepsy

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
· · ·	Nature Genetics, Volume 24, Number 4, April 2000, Escayg, A. et al, "Mutations of SCN1A, encoding a neuronal sodium channel, in two families	
	with GEFS+2", pages 343 to 345	1 - 4, 23, 24, 30 - 33,
X	See whole document	52
	AU 18465/01 A (McGILL UNIVERSITY) 4 June 2001	1 -11, 23, 24, 30 - 35
X	See page 58 line 12 to page 59 line 15, examples 3 and 6, and claims	52, 54 - 57, 60, 61, 65, 70, 73, 74
	Journal of Physiology, Volume 529, Number 3, 15 December 2000, Alekov,	
	A. K. et al, "A sodium channel mutation causing epilepsy in man exhibits	1 - 4, 23, 24, 30 - 33,
	subtle defects in fast inactivation and activation in vitro", pages 533 to 539	52
X	See Fig 1A, Abstract part 1.	

X Further documents are listed in the continuation of Box C X See patent family annex

*	Special categories of cited documents:	"T"	later document published after the international filing date or
"A"	document defining the general state of the art which is not considered to be of particular relevance		priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"E"	earlier application or patent but published on or after the international filing date	"X"	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an
"L"	document which may throw doubts on priority claim(s) or which is cited to establish the publication date of	"Y"	inventive step when the document is taken alone document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is
"O"	another citation or other special reason (as specified) document referring to an oral disclosure, use, exhibition or other means		combined with one or more other such documents, such combination being obvious to a person skilled in the art
"P"	document published prior to the international filing date but later than the priority date claimed	"&"	document member of the same patent family

Date of the actual completion of the international search

4 March 2002

Name and mailing address of the ISA/AU

AUSTRALIAN PATENT OFFICE

Date of mailing of the international search report

Authorized officer

PO BOX 200, WODEN ACT 2606, AUSTRALIA
E-mail address: pct@ipaustralia.gov.au
Facsimilc No. (02) 6285 3929

GAVIN THOMPSON
Telephone No : (02) 6283 2240

INTERNATIONAL SEARCH REPORT

International application No.

PCT/AU01/01648

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT						
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.				
P, X	Am. J. Hum. Genet., Volume 68, Number 4, April 2001, Wallace, R. H. et al, "Neuronal sodium-channel alpha1-subunit mutations in generalized epilepsy with febrile seizures plus", pages 859-865 See entire document	1 - 75				
	•					
	·					

INTERNATIONAL SEARCH REPORT

International application No. PCT/AU01/01648

Information on patent family members

This Annex lists the known "A" publication level patent family members relating to the patent documents cited in the above-mentioned international search report. The Australian Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Pater	nt Document Cited in Search Report			Patent Family Member	
AU	18465/01	WO	01/38564		
-					END OF ANNEX